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FILE 'USPATFULL' ENTERED AT 14:54:25 ON 27 APR 2011
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=> s (fluorescen? (3a) dye) (P) (masking (3a) dye)
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
FIELD CODE - 'AND' OPERATOR ASSUMED 'DYE) (P) '
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH

FIELD CODE - 'AND' OPERATOR ASSUMED 'DYE) (P) 'L1 122 (FLUORESCEN? (3A) DYE) (P) (MASKING (3A) DYE)

=> dup rem 11

PROCESSING COMPLETED FOR L1

L2 122 DUP REM L1 (0 DUPLICATES REMOVED)

=> d 12 1-20 ti

L2 ANSWER 1 OF 122 USPATFULL on STN

TI BGL4 Beta-Glucosidase and Nucleic Acids Encoding the Same

L2 ANSWER 2 OF 122 USPATFULL on STN

TI Method and Test Kit for the Rapid Identification and Characterization of Cells

L2 ANSWER 3 OF 122 USPATFULL on STN

- TI Protease Variants Active Over A Broad Temperature Range ANSWER 4 OF 122 USPATFULL on STN
- Surface Active Bleach and Dynamic pH
- ANSWER 5 OF 122 USPATFULL on STN
- Enzyme for the Production of Long Chain Peracid
- ANSWER 6 OF 122 USPATFULL on STN
- TI DUAL MECHANISM INHIBITORS FOR THE TREATMENT OF DISEASE
- L2 ANSWER 7 OF 122 USPATFULL on STN
- TI ENDOGLUCANASES
- ANSWER 8 OF 122 USPATFULL on STN
- ΤТ Novel Lipolytic Enzyme ELIP
- ANSWER 9 OF 122 USPATFULL on STN
- TI Methods for Improving Multiple Protein Properties
- ANSWER 10 OF 122 USPATFULL on STN
- Cleaning Enzymes and Malodor Prevention
- ANSWER 11 OF 122 USPATFULL on STN
- ΤI Modified Endoglucanase II and Methods of Use
- ANSWER 12 OF 122 USPATFULL on STN
- ΤI COMPOSITIONS AND METHODS COMPRISING SERINE PROTEASE VARIANTS
- ANSWER 13 OF 122 USPATFULL on STN
- USE OF PROTEIN HYDROLYSATES TO STABILIZE METALLOPROTEASE DETERGENT ΤI FORMULATIONS
- ANSWER 14 OF 122 USPATFULL on STN
- ΤI Stable Enzymatic Peracid Generating Systems
- ANSWER 15 OF 122 USPATFULL on STN
- TI BGL6 Beta-Glucosidase and Nucleic Acids Encoding the Same
- ANSWER 16 OF 122 USPATFULL on STN
- Novel Lipolytic Enzyme LIP2
- ANSWER 17 OF 122 USPATFULL on STN
- ΤI COMPOSITIONS AND METHODS COMPRISING A SUBTILISIN VARIANT
- ANSWER 18 OF 122 USPATFULL on STN
- TT COMPOSITIONS AND METHODS COMPRISING A SUBTILISIN VARIANT
- ANSWER 19 OF 122 USPATFULL on STN
- ΤI Cleaning Enzymes and Fragrance Production
- ANSWER 20 OF 122 USPATFULL on STN
- Cleaning Compositions Comprising Alpha-Galactosidase
- => d 11 21-122 ti
- ANSWER 21 OF 122 USPATFULL on STN
- TT Stable Enzymatic Peracid Generating Systems
- L1 ANSWER 22 OF 122 USPATFULL on STN

TI BGL6 Beta-Glucosidase and Nucleic Acids Encoding the Same ANSWER 23 OF 122 USPATFULL on STN TI Novel Lipolytic Enzyme LIP2 ANSWER 24 OF 122 USPATFULL on STN COMPOSITIONS AND METHODS COMPRISING A SUBTILISIN VARIANT ANSWER 25 OF 122 USPATFULL on STN TI COMPOSITIONS AND METHODS COMPRISING A SUBTILISIN VARIANT ANSWER 26 OF 122 USPATFULL on STN L1 TΙ Cleaning Enzymes and Fragrance Production ANSWER 27 OF 122 USPATFULL on STN ΤТ Cleaning Compositions Comprising Alpha-Galactosidase ANSWER 28 OF 122 USPATFULL on STN Novel lipolytic Enzyme lip1 TI ANSWER 29 OF 122 USPATFULL on STN Non-Phosphate Dish Detergents ANSWER 30 OF 122 USPATFULL on STN TI NOVEL BACILLUS 029cel CELLULASE ANSWER 31 OF 122 USPATFULL on STN Compositions and Methods Comprising Cellulase Variants with Reduced Affinity to Non-Cellulosic Materials ANSWER 32 OF 122 USPATFULL on STN TΙ ACYL Transferase Useful for Decontamination ANSWER 33 OF 122 USPATFULL on STN ΤI Novel Fungal Enzymes ANSWER 34 OF 122 USPATFULL on STN TΙ Novel Lipolytic Enzyme ELIP ANSWER 35 OF 122 USPATFULL on STN Thermostable Neutral Metalloproteases ANSWER 36 OF 122 USPATFULL on STN ΤI Perhydrolase Epitopes ANSWER 37 OF 122 USPATFULL on STN TT Variant humicola Grisea CBH1.1 ANSWER 38 OF 122 USPATFULL on STN ΤI BGL7 Beta-Glucosidase and Nucleic Acids Encoding The Same ANSWER 39 OF 122 USPATFULL on STN EGVII Endoglucanase and Nucleic Acids Encoding The Same ANSWER 40 OF 122 USPATFULL on STN L1 ΤI Novel CBH1 Homologs And Variant CBH1 Cellulases ANSWER 41 OF 122 USPATFULL on STN TT Composition Comprising A Coupled Enzyme System

ANSWER 42 OF 122 USPATFULL on STN

- TI Novel Fungal Enzymes
- ANSWER 43 OF 122 USPATFULL on STN
- Bacillus mHKcel Cellulase
- ANSWER 44 OF 122 USPATFULL on STN
- Variant Humicola Grisea CBH1.1
- ANSWER 45 OF 122 USPATFULL on STN
- KAPPA-CARRAGEENASE AND KAPPA-CARRAGEENASE-CONTAINING COMPOSITIONS
- L1 ANSWER 46 OF 122 USPATFULL on STN
- TI OPTICALLY-DETECTABLE ENZYME SUBSTRATES AND THEIR METHOD OF USE
- ANSWER 47 OF 122 USPATFULL on STN
- ΤТ EGVI Endoglucanase and Nucleic Acids Encoding the Same
- TI IN VIVO OPTICAL IMAGING
- ANSWER 49 OF 122 USPATFULL on STN

ANSWER 48 OF 122 USPATFULL on STN

- MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN THE OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS
- ANSWER 50 OF 122 USPATFULL on STN
- TΤ POLYGE OXIDASES
- ANSWER 51 OF 122 USPATFULL on STN
- Use and production of storage-stable neutral metalloprotease
- ANSWER 52 OF 122 USPATFULL on STN L1
- ΤI Compositions and methods utilizing DNA polymerases
- ANSWER 53 OF 122 USPATFULL on STN
- Cleaning compositions comprising transglucosidase
- ANSWER 54 OF 122 USPATFULL on STN
- TΙ Novel Laccases, Compositions And Methods of Use
- ANSWER 55 OF 122 USPATFULL on STN
- LACCASE MEDIATORS AND METHODS OF USE
- ANSWER 56 OF 122 USPATFULL on STN
- BGL5 Beta-glucosidase and nucleic acids encoding the same ΤI
- ANSWER 57 OF 122 USPATFULL on STN
- TT Perhydrolase
- ANSWER 58 OF 122 USPATFULL on STN
- TI BGL3 Beta-Glucosidase and nucleic acids encoding the same
- ANSWER 59 OF 122 USPATFULL on STN
- Protease variants active over a broad temperature range
- ANSWER 60 OF 122 USPATFULL on STN L1
- TI Surface active bleach and dynamic pH
- ANSWER 61 OF 122 USPATFULL on STN
- TT Detergents with stabilized enzyme systems
- L1 ANSWER 62 OF 122 USPATFULL on STN

- Novel Liplytic Enzyme Elip
- ANSWER 63 OF 122 USPATFULL on STN
- TI Novel Lipolytic Enzyme Lip2
- ANSWER 64 OF 122 USPATFULL on STN
- EGVI endoglucanase and nucleic acids encoding the same
- ANSWER 65 OF 122 USPATFULL on STN
- EGVII endoglucanase and nucleic acids encoding the same
- L1 ANSWER 66 OF 122 USPATFULL on STN
- TI Novel variant hypocrea jercorina CBH1 cellulases
- ANSWER 67 OF 122 USPATFULL on STN
- ΤТ Novel bacillus 029cel cellulase
- ANSWER 68 OF 122 USPATFULL on STN TI Enzyme for the production of long chain peracid
- ANSWER 69 OF 122 USPATFULL on STN
- Novel bacillus bagcel cellulase
- ANSWER 70 OF 122 USPATFULL on STN
- TI Novel bacillus mhkcel cellulase
- ANSWER 71 OF 122 USPATFULL on STN
- Novel EGIII-like enzymes, DNA encoding such enzymes and methods for producing such enzymes
- ANSWER 72 OF 122 USPATFULL on STN
- TΙ Bgl6 beta-glucosidase and nucleic acids encoding the same
- ANSWER 73 OF 122 USPATFULL on STN
- Novel variant hypocrea jecorina CBH2 cellulases
- ANSWER 74 OF 122 USPATFULL on STN
- EGVI endoglucanase and nucleic acids encoding the same
- ANSWER 75 OF 122 USPATFULL on STN EGVI endoglucanase and nucleic acids encoding the same
- ANSWER 76 OF 122 USPATFULL on STN
- ΤI EGVII endoglucanase and nucleic acids encoding the same
- ANSWER 77 OF 122 USPATFULL on STN TT
- EGVII endoglucanase and nucleic acids encoding the same
- TI Exo-endo cellulase fusion protein
- ANSWER 78 OF 122 USPATFULL on STN
- ANSWER 79 OF 122 USPATFULL on STN
- BGL4 beta-glucosidase and nucleic acids encoding the same
- ANSWER 80 OF 122 USPATFULL on STN L1
- ΤI Novel variant hypocrea jecorina CBHlcellulases
- ANSWER 81 OF 122 USPATFULL on STN
- TT BGL5 beta-glucosidase and nucleic acids encoding the same
- L1 ANSWER 82 OF 122 USPATFULL on STN

- TΤ Optically-detectable enzyme substrates and their method of use
- ANSWER 83 OF 122 USPATFULL on STN
- BGL3 beta-glucosidase and nucleic acids encoding the same
- ANSWER 84 OF 122 USPATFULL on STN
- BGL3 beta-glucosidase and nucleic acids encoding the same
- ANSWER 85 OF 122 USPATFULL on STN
- Natural product based apoptosis inducers
- L1 ANSWER 86 OF 122 USPATFULL on STN
- TI Novel CBH1 homologs and variant CBH1 cellulases
- ANSWER 87 OF 122 USPATFULL on STN
- ΤТ Variant humicola grisea CBH1.1
- ANSWER 88 OF 122 USPATFULL on STN
- TI
- BGL7 beta-glucosidase and nucleic acids encoding the same
- ANSWER 89 OF 122 USPATFULL on STN
- Compositions and methods utilizing DNA polymerases
- ANSWER 90 OF 122 USPATFULL on STN
- ΤI Mutant EGIII cellulase, DNA encoding such EGIII compositions and methods for obtaining same
- ANSWER 91 OF 122 USPATFULL on STN
- Image forming device ANSWER 92 OF 122 USPATFULL on STN L1
- CHRYSOSPORIUM CELLULASE AND METHODS OF USE TΙ
- ANSWER 93 OF 122 USPATFULL on STN
- Compositions and methods utilizing DNA polymerases
- ANSWER 94 OF 122 USPATFULL on STN
- TΙ BGL4 beta-glucosidase and nucleic acids encoding the same
- ANSWER 95 OF 122 USPATFULL on STN
- BGL5 beta-glucosidase and nucleic acids encoding the same
- ANSWER 96 OF 122 USPATFULL on STN
- ΤI EGVIII endoglucanase and nucleic acids encoding the same
- ANSWER 97 OF 122 USPATFULL on STN
- TT EGVII endoglucanase and nucleic acids encoding the same
- TI
- EGVI endoglucanase and nucleic acids encoding the same
- ANSWER 99 OF 122 USPATFULL on STN
- Variant EGIII-like cellulase compositions

ANSWER 98 OF 122 USPATFULL on STN

- ANSWER 100 OF 122 USPATFULL on STN L1
- ΤI Masking of the background fluorescence and luminescence in the optical analysis of biomedical assays
- ANSWER 101 OF 122 USPATFULL on STN
- ΤТ Mutant EGIII cellulase, DNA encoding such EGIII compositions and methods for obtaining same

- L1 ANSWER 102 OF 122 USPATFULL on STN
- TI BGL3 beta-glucosidase and nucleic acids encoding the same
- L1 ANSWER 103 OF 122 USPATFULL on STN
- TI Cellulase for use in industrial processes
- L1 ANSWER 104 OF 122 USPATFULL on STN
  - TI Mutant EGIII cellulase, DNA encoding such EGIII compositions and methods for obtaining same
- L1 ANSWER 105 OF 122 USPATFULL on STN
- TI Method and compositions for treating cellulose containing fabrics using truncated cellulase enzyme compositions
- L1 ANSWER 106 OF 122 USPATFULL on STN
- TI Cellulase for use in industrial processes
- L1 ANSWER 107 OF 122 USPATFULL on STN
- TI Mutant EGIII cellulase, DNA encoding such EGIII compositions and methods for obtaining same
- L1 ANSWER 108 OF 122 USPATFULL on STN
- TI MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS
- L1 ANSWER 109 OF 122 USPATFULL on STN
- TI Masking background fluorescence and luminescence in optical analysis of biomedical assays
- L1 ANSWER 110 OF 122 USPATFULL on STN
- TI Masking of the background fluorescence and luminescence in the optical analysis of biomedical assays
- L1 ANSWER 111 OF 122 USPATFULL on STN
- TI Variant EGIII-like cellulase compositions
- L1 ANSWER 112 OF 122 USPATFULL on STN
- TI Method and compositions for treating cellulose containing fabrics using truncated cellulase enzyme compositions
- L1 ANSWER 113 OF 122 USPATFULL on STN
- TI Mutant EGIII cellulase, DNA encoding such EGIII compositions and methods for obtaining same
- L1 ANSWER 114 OF 122 USPATFULL on STN
- TI Purified cellulase and method of producing
- L1 ANSWER 115 OF 122 USPATFULL on STN
- TI Treating cellulosic materials with cellulases from chrysosporium
- L1 ANSWER 116 OF 122 USPATFULL on STN
- TI Oversized cellulase compositions for use in detergent compositions and in the treatment of textiles
- L1 ANSWER 117 OF 122 USPATFULL on STN
- TI Mutant Thermonospora spp. cellulase
- L1 ANSWER 118 OF 122 USPATFULL on STN
- TI Cellulase compositions and methods of use

- L1 ANSWER 119 OF 122 USPATFULL on STN
- TT Chromene dves
- L1 ANSWER 120 OF 122 USPATFULL on STN
- TI Article identification material and method and apparatus for using it
- ANSWER 121 OF 122 USPATFULL on STN
- INSPECTION PENETRANT DEVELOPMENT PROCESS EMPLOYING FUSIBLE WAXES
- L1 ANSWER 122 OF 122 USPATFULL on STN
- DEVELOPERS FOR INSPECTION PENETRANTS EMPLOYING FUSIBLE WAXES TI

#### => d 11 100-122 ibib abs

L1 ANSWER 100 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2003:133996 USPATFULL <<LOGINID::20110427>>

Masking of the background fluorescence and luminescence TITLE:

in the optical analysis of biomedical assays INVENTOR(S): Krahn, Thomas, Hagen, GERMANY, FEDERAL REPUBLIC OF

Paffhausen, Wolfgang, Leverkusen, GERMANY, FEDERAL

REPUBLIC OF

Schade, Andreas, Essen, GERMANY, FEDERAL REPUBLIC OF Bechem, Martin, Wuppertal, GERMANY, FEDERAL REPUBLIC OF Schmidt, Delf, Wuppertal, GERMANY, FEDERAL REPUBLIC OF

NUMBER KIND DATE US 20030092081 A1 20030515 US 7138280 B2 20061121 US 2002-263607 A1 20021003 (10) PATENT INFORMATION:

APPLICATION INFO.:

Division of Ser. No. US 2001-966522, filed on 28 Sep RELATED APPLN. INFO.: 2001, PENDING

NUMBER DATE PRIORITY INFORMATION: DE 1996-19621312 19960528

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: KURT BRISCOE, NORRIS, MCLAUGHLIN & MARCUS, P.A., 220 EAST 42ND STREET, 30TH FLOOR, NEW YORK, NY, 10017

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: 10 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 438

CAS INDEXING IS AVAILABLE FOR THIS PATENT. AB

In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3

containing the fluorescent dye 4. The sensitivity of analytical detection can be considerably improved if to the

fluorescent dye 4 already present in addition a

masking dve 9, which absorbs the excitation light 6 for the fluorescent dve 4 and/or its emission light

7, is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6 or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating layer 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously,

these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor layer 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here if a masking dye 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission light or (in the case of luminescent ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye in the solution 3 or optionally as an additional measure, a separating layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 101 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2003:124326 USPATFULL <<LOGINID::20110427>>

TITLE: Mutant EGIII cellulase, DNA encoding such EGIII compositions and methods for obtaining same

INVENTOR(S): Fowler, Timothy, Bainbridge Island, UNITED KINGDOM
Mitchinson, Colin, Half Moon Bay, CA, UNITED STATES

can be applied to the cell or receptor layer 12 at the bottom 2.

NUMBER KIND DATE

PATENT INFORMATION: US 20030084515 Al 20030508
US 6582750 B2 20030624
APPLICATION INFO: US 2002-261997 Al 20020930 (10)

RELATED APPLN. INFO.: Division of Ser. No. US 2002-75872, filed on 13 Feb. 2002, GRANTED, Pat. No. US 5500211 Continuation of Ser. No. US 2000-633084, filed on 4 Aug 2000, GRANTED, Pat. No. US 2000-633084, filed on 4 Aug 2000, GRANTED, Pat.

No. US 6407046 Continuation-in-part of Ser. No. US 1998-146770, filed on 3 Sep 1998, GRANTED, Pat. No. US 6187732

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Genencor International, Inc., 925 Page Mill Road, Palo

Alto, CA, 94034-1013

NUMBER OF CLAIMS: 17 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 5 Drawing Page(s)
LINE COUNT: 1685

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention relates to variant EGIII cellulases that have improved stability and/or performance. The variant cellulases have replacements at sensitive residues to improve stability and/or

performance.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 102 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2003:120298 USPATFULL <<LOGINID::20110427>>

TITLE: BGL3 beta-glucosidase and nucleic acids encoding the

same

INVENTOR(S): Dunn-Coleman, Nigel, Los Gatos, CA, UNITED STATES
Geodegebuur, Frits, Vlaardingen, NETHERLANDS

Geodegebuur, Frits, Vlaardingen, NETHERLANDS Ward, Michael, San Francisco, CA, UNITED STATES

Yao, Jian, Sunnyvale, CA, UNITED STATES

NUMBER KIND DATE PATENT INFORMATION: US 20030082779 A1 20030501 US 6982159 B2 20060103 US 2001-957880 A1 20010921 (9) APPLICATION INFO.: DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: VICTORIA L. BOYD, GENENCOR INTERNATIONAL, INC., 925

PAGE MILL ROAD, PALO ALTO, CA, 94034-1013

NUMBER OF CLAIMS: 28

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 2 Drawing Page(s) 1915

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides a novel β-glucosidase nucleic acid

sequence, designated bgl3, and the corresponding BGL3 amino acid sequence. The invention also provides expression vectors and host cells comprising a nucleic acid sequence encoding BGL3, recombinant BGL3

proteins and methods for producing the same.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 103 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2002:337907 USPATFULL <<LOGINID::20110427>>

TITLE: Cellulase for use in industrial processes

INVENTOR(S): Clarkson, Kathleen A., San Francisco, CA, UNITED STATES Swanson, Barbara, San Francisco, CA, UNITED STATES

Winetzky, Deborah, South San Francisco, CA, UNITED

STATES

NUMBER KIND DATE PATENT INFORMATION: US 20020193272 A1 20021219 APPLICATION INFO.: US 2002-172480 A1 20020614 (10)

RELATED APPLN. INFO.: Division of Ser. No. US 1996-719506, filed on 25 Sep

1996, PENDING

DOCUMENT TYPE: Utility FILE SEGMENT:

APPLICATION LEGAL REPRESENTATIVE: Genencor International, Inc., 925 Page Mill Road, Palo

Alto, CA, 94034-1013

NUMBER OF CLAIMS: 19

EXEMPLARY CLAIM: NUMBER OF DRAWINGS: 6 Drawing Page(s) LINE COUNT: 1224

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A method for treating cellulosic materials is disclosed which comprises contacting the cellulosic material with a cellulase obtainable from Thermomonospora fusca corresponding to E5 or a derivative thereof. Particularly preferred methods comprise stonewashing and detergent cleaning of cotton fabrics, the production of paper products, as an additive to animal feed and in the production of food, starch, ethanol and sugar.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 104 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2002:295069 USPATFULL <<LOGINID::20110427>>

TITLE: Mutant EGIII cellulase, DNA encoding such EGIII compositions and methods for obtaining same

INVENTOR(S): Fowler, Timothy, Bainbridge Island, WA, UNITED STATES

Mitchinson, Colin, Half Moon Bay, CA, UNITED STATES

RELATED APPLN. INFO:: Continuation of Ser. No. US 2000-633084, filed on 4 Aug 2000, PENDING Continuation-in-part of Ser. No. US

1998-146770, filed on 3 Sep 1998, GRANTED, Pat. No. US 6187732

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Genencor International, Inc., 925 Page Mill Road, Palo Alto, CA, 94034-1013

NUMBER OF CLAIMS: 17 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 5 Drawing Page(s)

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention relates to variant EGIII cellulases that have improved stability and/or performance. The variant cellulases have replacements at sensitive residues to improve stability and/or performance.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 105 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2002:294731 USPATFULL <<LOGINID::20110427>>

TITLE: Method and compositions for treating cellulose containing fabrics using truncated cellulase enzyme

compositions
INVENTOR(S): Fowler, Timothy Fowler, San Carlos, CA, UNITED STATES

Clarkson, Kathleen A., San Francisco, CA, UNITED STATES

Ward, Michael, San Francisco, CA, UNITED STATES Collier, Katherine D., Redwood City, CA, UNITED STATES

Larenas, Edmund, Moss Beach, CA, UNITED STATES

NUMBER KIND DATE

PATENT INFORMATION: US 20020164774 A1 20021107
US 6620605 B2 20030916

APPLICATION INFO: US 2001-916494 A1 20010727 (9)

RELATED APPLN. INFO.: Continuation of Ser. No. US 1995-382452, filed on 1 Feb

1993-169948, filed on 17 Dec 1993, PATENTED

DOCUMENT TYPE: Utility PATENTED

FILE SEGMENT: APPLICATION

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Genencor International, Inc., 925 Page Mill Road, Palo Alto, CA, 94034-1013

NUMBER OF CLAIMS: 38

EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 13 Drawing Page(s)

LINE COUNT: 2890

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

B Improved methods of treating cellulose containing fabrics with cellulase comprising contacting the cellulose fabrics with truncated cellulase enzyme. Treatment of cellulose containing fabrics with cellulase core domains of the invention are disclosed as offering specific advantages of reduced redeposition of dye and increased abrasion.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 106 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2002:238459 USPATFULL <<LOGINID::20110427>>

TITLE: Cellulase for use in industrial processes

INVENTOR(S): Clarkson, Kathleen A., San Francisco, CA, United States Swanson, Barbara, San Francisco, CA, United States Winetzky, Deborah, South San Francisco, CA, United

States

PATENT ASSIGNEE(S): Genencor International, Inc., Palo Alto, CA, United

States (U.S. corporation)

NUMBER KIND DATE US 6451063 B1 20020917 US 1996-719506 19960925 PATENT INFORMATION: APPLICATION INFO.: 19960925 (8)

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

PRIMARY EXAMINER: Diamond, Alan

LEGAL REPRESENTATIVE: Genencor International, Inc NUMBER OF CLAIMS: 11

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 11 D: 1226 11 Drawing Figure(s); 6 Drawing Page(s)

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A method for treating cellulosic materials is disclosed which comprises contacting the cellulosic material with a cellulase obtainable from Thermomonospora fusca corresponding to E5 or a derivative thereof. Particularly preferred methods comprise stonewashing and detergent cleaning of cotton fabrics, the production of paper products, as an additive to animal feed and in the production of food, starch, ethanol and sugar.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 107 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2002:144226 USPATFULL <<LOGINID::20110427>> Mutant EGIII cellulase, DNA encoding such EGIII TITLE:

compositions and methods for obtaining same

INVENTOR(S): Fowler, Timothy, Bainbridge Island, WA, United States Mitchinson, Colin, Half Moon Bay, CA, United States

Genencor International, Inc., Palo Alto, CA, United PATENT ASSIGNEE(S):

States (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: US 6407046 B1 20020618 APPLICATION INFO.: US 2000-633084 20000804 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1998-146770, filed on 3 Sep 1998, now patented, Pat. No. US 6187732

DOCUMENT TYPE: Utility

FILE SEGMENT: GRANTED

FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Delcotto, Gregory
LEGAL REPRESENTATIVE: Genencor International, Inc.

NUMBER OF CLAIMS: 12 12

NUMBER OF DRAWINGS: 5 Drawing Figure(s); 5 Drawing Page(s)

1627

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to variant EGIII cellulases that have improved stability and/or performance. The variant cellulases have replacements at sensitive residues to improve stability and/or

performance.

### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 108 OF 122 USPATFULL on STN

ACCESSION NUMBER:

2002:37557 USPATFULL <<LOGINID::20110427>>

TITLE: MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS

KRAHN, THOMAS, HAGEN, GERMANY, FEDERAL REPUBLIC OF INVENTOR(S): PAFFHAUSEN, WOLFGANG, LEVERKUSEN, GERMANY, FEDERAL

REPUBLIC OF

SCHADE, ANDREAS, ESSEN, GERMANY, FEDERAL REPUBLIC OF BECHEM, MARTIN, WUPPERTAL, GERMANY, FEDERAL REPUBLIC OF SCHMIDT, DELF, WUPPERTAL, GERMANY, FEDERAL REPUBLIC OF

NUMBER KIND DATE US 20020022274 A1 20020221 US 6420183 B2 20020716 US 1998-194099 A1 19981120 (9) WO 1997-EP2662 19970523 PATENT INFORMATION: APPLICATION INFO.: NUMBER DATE

PRIORITY INFORMATION: DE 1996-19621312 19960528

DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

MORRIS MCLAUGHLIN & MARCUS, P.A 30TH FLOOR, NEW YORK, NY, 10017 6 LEGAL REPRESENTATIVE: NORRIS McLAUGHLIN & MARCUS, P.A., 220 EAST 42nd STREET

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 10 Drawing Page(s) LINE COUNT:

462 CAS INDEXING IS AVAILABLE FOR THIS PATENT.

In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3

containing the fluorescent dye 4. The sensitivity of

analytical detection can be considerably improved if to the fluorescent dye 4 already present in addition a

masking dve 9, which absorbs the excitation light 6

for the fluorescent dve 4 and/or its emission light 7. is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6

or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating laver 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously, these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor layer 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which

a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here

if a masking dye 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission

light or (in the case of luminescent ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye

in the solution 3 or optionally as an additional measure, a separating

layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light can be applied to the cell or receptor layer 12 at the bottom 2.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 109 OF 122 USPATFULL on STN

ACCESSION NUMBER:

2002:27123 USPATFULL <<LOGINID::20110427>> TITLE: Masking background fluorescence and luminescence in optical analysis of biomedical assays

INVENTOR(S): Krahn, Thomas, Hagen, GERMANY, FEDERAL REPUBLIC OF Paffhausen, Wolfgang, Leverkusen, GERMANY, FEDERAL REPUBLIC OF Schade, Andreas, Essen, GERMANY, FEDERAL REPUBLIC OF

Bechem, Martin, Wuppertal, GERMANY, FEDERAL REPUBLIC OF Schmidt, Delf, Wuppertal, GERMANY, FEDERAL REPUBLIC OF

NUMBER KIND DATE US 20020015969 A1 20020207 US 7063952 B2 20060620 US 2001-966137 A1 20010928 PATENT INFORMATION: APPLICATION INFO.:

RELATED APPLN. INFO.: Division of Ser. No. US 1998-194099, filed on 20 Nov 1998, PENDING

NUMBER DATE
----DE 1996-19621312 19960528 PRIORITY INFORMATION: DOCUMENT TYPE: Utility

FILE SEGMENT:

APPLICATION LEGAL REPRESENTATIVE: Kurt G. Briscoe, Norris McLaughlin & Marcus, P.A., 30th Floor, 220 East 42nd Street, New York, NY, 10017

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

10 Drawing Page(s) NUMBER OF DRAWINGS: LINE COUNT: 462

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3 containing the fluorescent dve 4. The sensitivity of analytical detection can be considerably improved if to the fluorescent dve 4 already present in addition a masking dye 9, which absorbs the excitation light 6 for the fluorescent dye 4 and/or its emission light 7, is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6 or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating layer 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously, these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor layer 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here

if a masking dye 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission light or (in the case of luminescent ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye in the solution 3 or optionally as an additional measure, a separating layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light can be applied to the cell or receptor layer 12 at the bottom 2.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 110 OF 122 USPATFULL on STN ACCESSION NUMBER:

TITLE:

2002:16874 USPATFULL <<LOGINID::20110427>> Masking of the background fluorescence and luminescence in the optical analysis of biomedical assays

INVENTOR(S):

Krahn, Thoams, Hagen, GERMANY, FEDERAL REPUBLIC OF Paffhausen, Wolfgang, Leverkusen, GERMANY, FEDERAL REPUBLIC OF

Schade, Andreas, Essen, GERMANY, FEDERAL REPUBLIC OF Bechem, Martin, Wuppertal, GERMANY, FEDERAL REPUBLIC OF Schmidt, Delf, Wuppertal, GERMANY, FEDERAL REPUBLIC OF

NUMBER KIND DATE US 20020009754 A1 20020124 US 2001-966522 A1 20010928 (9)

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

Continuation of Ser. No. US 1998-194099, filed on 20 Nov 1998, PENDING

NUMBER DATE NUMBER DATE
DE 1996-19621312 19960528

PRIORITY INFORMATION: DOCUMENT TYPE:

Utility

FILE SEGMENT: APPLICATION LEGAL REPRESENTATIVE: Kurt G. Briscoe, Norris McLaughlin & Marcus, P.A., 30th

Floor, 220 East 42nd Street, New York, NY, 10017

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 10 Drawing Page(s) LINE COUNT: 462

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3 containing the fluorescent dve 4. The sensitivity of analytical detection can be considerably improved if to the fluorescent dye 4 already present in addition a masking dye 9, which absorbs the excitation light 6 for the fluorescent dye 4 and/or its emission light

7, is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6 or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating layer 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously, these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor layer 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here

if a masking dye 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission light or (in the case of luminescent ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye in the solution 3 or optionally as an additional measure, a separating layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light can be applied to the cell or receptor layer 12 at the bottom 2. (FIGS. 2 and 10)

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 111 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2001:121435 USPATFULL <<LOGINID::20110427>> TITLE: Variant EGIII-like cellulase compositions

INVENTOR(S): Mitchinson, Colin, Half Moon Bay, CA, United States Wendt, Dan J., Walnut Creek, CA, United States

Genencor International, Inc., Palo Alto, CA, United PATENT ASSIGNEE(S):

States (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: US 6268328 B1 20010731 APPLICATION INFO:: US 1998-216295 19981218 APPLICATION INFO.: 19981218 (9) DOCUMENT TYPE: Utility GRANTED FILE SEGMENT:

FILE SEGMENT: GRANIED

PRIMARY EXAMINER: Gupta, Yogendra N.

ASSISTANT EXAMINER: Elhilo, Eisa

LEGAL REPRESENTATIVE: Genencor International, Inc.

NUMBER OF CLAIMS: 20 EXEMPLARY CLAIM:

7 Drawing Figure(s); 7 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 1619

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to novel variant EGIII or EGIII-like cellulases which have improved stability. The variant cellulases have performance sensitive residues replaced to a residue having improved stability.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 112 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2001:121303 USPATFULL <<LOGINID::20110427>> TITLE: Method and compositions for treating cellulose

containing fabrics using truncated cellulase enzyme

compositions

Fowler, Timothy, San Carlos, CA, United States INVENTOR(S): Clarkson, Kathleen A., San Francisco, CA, United States Ward, Michael, San Francisco, CA, United States

Collier, Katherine D., Redwood City, CA, United States Larenas, Edmund, Moss Beach, CA, United States

PATENT ASSIGNEE(S): Genencor International, Inc., Rochester, NY, United

States (U.S. corporation)

NUMBER KIND DATE US 6268196 B1 20010731 US 1995-382452 19950201 (8) PATENT INFORMATION: APPLICATION INFO.:

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1993-169948, filed

on 17 Dec 1993

DOCUMENT TYPE: Utility

FILE SEGMENT: GRANTED
PRIMARY EXAMINER: Patterson, Jr., Charles L.

LEGAL REPRESENTATIVE: Marcus-Werner, LynnGenecor International, Inc.

NUMBER OF CLAIMS: 48 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 31 Drawing Figure(s); 24 Drawing Page(s)

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Improved methods of treating cellulose containing fabrics with cellulase comprising contacting the cellulose fabrics with truncated cellulase enzyme. Treatment of cellulose containing fabrics with cellulase core domains of the invention are disclosed as offering specific advantages

of reduced redeposition of dye and increased abrasion.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 113 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2001:22178 USPATFULL <<LOGINID::20110427>>

TITLE: Mutant EGIII cellulase, DNA encoding such EGIII compositions and methods for obtaining same

INVENTOR(S): Fowler, Timothy, Bainbridge Island, WA, United States Mitchinson, Colin, Half Moon Bay, CA, United States

Genencor International, Inc., Rochester, NY, United PATENT ASSIGNEE(S):

States (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: US 6187732 B1 20010213 US 1998-146770 19980903 (9) APPLICATION INFO.: DOCUMENT TYPE: Utility

Granted FILE SEGMENT: PRIMARY EXAMINER: Fries, Kery

LEGAL REPRESENTATIVE: Faris, Susan K.Genencor International, Incorporated

NUMBER OF CLAIMS: 10 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 4 Drawing Figure(s); 3 Drawing Page(s)

LINE COUNT: 1222

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to variant EGIII cellulases which have improved stability and/or performance. The variant cellulases have replacements at sensitive residues to improve stability and/or

performance.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 114 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2000:9859 USPATFULL <<LOGINID::20110427>> TITLE: Purified cellulase and method of producing

INVENTOR(S): Bower, Benjamin S., Pacifica, CA, United States Clarkson, Kathleen A., San Francisco, CA, United States

Collier, Katherine D., Redwood City, CA, United States Kellis, James T., Portola Valley, CA, United States Kelly, Moira B., San Francisco, CA, United States Larenas, Edmund A., Moss Beach, CA, United States

PATENT ASSIGNEE(S): Genencor International, Inc., Rochester, NY, United States (U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 6017870 20000125 APPLICATION INFO.: US 1996-728350 19961009 (8)

DOCUMENT TYPE: Utility FILE SEGMENT: Granted
PRIMARY EXAMINER: Fries, Kery

LEGAL REPRESENTATIVE: Stone, Christopher L.Genencor International, Inc.

NUMBER OF CLAIMS: 19 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 3 Drawing Figure(s); 3 Drawing Page(s) LINE COUNT: 891

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A purified novel cellulase composition is provided which may be isolated from a fermentation culture of Trichoderma longibrachiatum and has a molecular weight of about 95-105 kD as approximated on SDS-PAGE (see FIG. 1), a pl of about 5.6-6.8 as estimated on an IEF gel and a pH optimum of about 5.0 on RBB-CMC when measured at 65° C. and pH 4 or lower at temperatures of 40° C. and 50° C.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 115 OF 122 USPATFULL on STN

ACCESSION NUMBER: 2000:7208 USPATFULL <<LOGINID::20110427>>

TITLE: Treating cellulosic materials with cellulases from

chrysosporium INVENTOR(S):

Emalfarb, Mark Aaron, Jupiter, FL, United States Ben-Bassat, Arie, Wilmington, DE, United States Sinitsyn, Arkady Panteleimonovich, Moscow, Russian

Federation Emalfarb, Mark A., Jupiter, FL, United States (U.S. PATENT ASSIGNEE(S):

individual)

NUMBER KIND DATE PATENT INFORMATION: US 6015707 APPLICATION INFO.: US 1998-106026 20000118 19980629 (9)

RELATED APPLN. INFO.: Division of Ser. No. US 1996-731170, filed on 10 Oct

1996, now patented, Pat. No. US 5811381

DOCUMENT TYPE: Utility FILE SEGMENT: Granted PRIMARY EXAMINER: Wax, Robert A.

LEGAL REPRESENTATIVE: Morgan & Finnegan, LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM: LINE COUNT: 1900

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The subject invention relates to novel compositions of neutral and/or alkaline cellulase and methods for obtaining neutral and/or alkaline cellulase compositions from Chrysosporium cultures, in particular Chrysosporium lucknowense. This invention also provides mutants and methods of generating mutants of Chrysosporium capable of producing neutral and/or alkaline cellulase. This invention also relates to the genes encoding the enzymes comprising the neutral and/or alkaline cellulase composition. In addition, this invention provides methods of culturing Chrysosporium to produce neutral and/or alkaline cellulases. The neutral and/or alkaline cellulase compositions of the subject invention can be used in a variety of processes including stone washing of clothing, detergent processes, deinking and biobleaching of paper & pulp and treatment of waste streams.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 116 OF 122 USPATFULL on STN

ACCESSION NUMBER: 1999:151002 USPATFULL <<LOGINID::20110427>>

TITLE: Oversized cellulase compositions for use in detergent

compositions and in the treatment of textiles

Bower, Benjamin S., Pacifica, CA, United States INVENTOR(S):

Clarkson, Kathleen A., San Francisco, CA, United States

Larenas, Edmund A., Moss Beach, CA, United States Ward, Michael, San Francisco, CA, United States

Genencor International, Inc., Rochester, NY, United PATENT ASSIGNEE(S): States (U.S. corporation)

NUMBER KIND DATE US 5989899 19991123 US 1996-774065 19961223 (8) PATENT INFORMATION: APPLICATION INFO.: DOCUMENT TYPE: Utility FILE SEGMENT: Granted PRIMARY EXAMINER: Patterson, Jr., Charles

LEGAL REPRESENTATIVE: Stone, Christopher L. NUMBER OF CLAIMS: 10

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 11 Drawing Figure(s); 9 Drawing Page(s) LINE COUNT: 1212

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to a modified cellulase protein which is advantageously used in the treatment of textiles. Particularly, a method for treating a cellulose containing fabric is provided comprising the steps of forming an aqueous solution comprising a cellulase composition which differs from a precursor cellulase in that it has been enlarged and contacting the aqueous solution with a cellulose containing fabric for a time and under conditions appropriate to treat the fabric. The enlarged cellulase may comprise a multimeric composition of two or more distinct cellulase units or a single cellulase which has had adhered thereto polymeric or fibrous constituents.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 117 OF 122 USPATFULL on STN

ACCESSION NUMBER: 1999:21543 USPATFULL <<LOGINID::20110427>>

TITLE: Mutant Thermonospora spp. cellulase

INVENTOR(S): Goedegebuur, Frits, Vloordingen, Netherlands Power, Scott D., San Bruno, CA, United States

Winetzky, Deborah, Foster City, CA, United States Van Kimmenade, Anita, San Bruno, CA, United States Yoon, Mee-Young, Palo Alto, CA, United States

PATENT ASSIGNEE(S): Genencor International, Inc., Rochester, NY, United States (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: US 5871550 19990216 APPLICATION INFO.: US 1997-924440 19970826 (8) DOCUMENT TYPE: Utility

DOCUMENT TYPE: FILE SEGMENT: FILE SEGMENT: Granted
PRIMARY EXAMINER: Fries, Kery
LEGAL REPRESENTATIVE: Stone, Christopher L.
NUMBER OF CLAIMS: 11

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 5 Drawing Figure(s); 4 Drawing Page(s) LINE COUNT: 1297

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A mutant cellulase obtainable from Thermomonospora spp is provided which differs from a precursor cellulase in that it has been genetically

engineered to introduce a substitution, deletion or addition of an amino acid residue to said precursor cellulase which provided improved activity in a detergent. Preferably, the substitution is at a residue corresponding to Tild in Thermomonospora fusca.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 118 OF 122 USPATFULL on STN

ACCESSION NUMBER: 1998:115696 USPATFULL <<LOGINID::20110427>>

TITLE: Cellulase compositions and methods of use INVENTOR(S): Emalfarb, Mark Aaron, Jupiter, FL, United

NVENTOR(S): Emalfarb, Mark Aaron, Jupiter, FL, United States Ben-Bassat, Arie, Wilmington, DE, United States Burlingame, Richard P., Manitowoc, WI, United States Chernoglazov, Vladimir Mikhaylovich, Moscow, Russian

Federation

Okounev, Oleg Nicolaevich, Moscow, Russian Federation Olson, Philip T., Manitowoc, WI, United States Sinitsyn, Arkady Panteleimonovich, Moscow, Russian

Federation Federation

Solovjeva, Irina Vladimirovna, Moscow Region, Russian Federation

PATENT ASSIGNEE(S): Emalfarb, Mark A., Jupiter, FL, United States (U.S.

individual)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Lau, Kawai

LEGAL REPRESENTATIVE: Morgan & Finnegan

NUMBER OF CLAIMS: 44
EXEMPLARY CLAIM: 12
LINE COUNT: 2026

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The subject invention relates to novel compositions of neutral and/or alkaline cellulase and methods for obtaining neutral and/or alkaline cellulase compositions from Chrysosporium cultures, in particular Chrysosporium lucknowense. This invention also provides mutants and methods of generating mutants of Chrysosporium capable of producing neutral and/or alkaline cellulose. This invention also relates to the genes encoding the enzymes comprising the neutral and/or alkaline cellulase composition. In addition, this invention provides methods of culturing Chrysosporium to produce neutral and/or alkaline cellulases. The neutral and/or alkaline cellulase compositions of the subject invention can be used in a variety of processes including stone washing of clothing, detergent processes, deinking and biobleaching of paper & pulp and treatment of waste streams.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 119 OF 122 USPATFULL on STN

ACCESSION NUMBER: 1998:14946 USPATFULL <<LOGINID::20110427>>

TITLE: Chromene dyes

INVENTOR(S): DeBoer, Charles David, Palmyra, NY, United States Robello, Douglas Robert, Webster, NY, United States

Tutt, Lee William, Webster, NY, United States

PATENT ASSIGNEE(S): Eastman Kodak Company, Rochester, NY, United States

(U.S. corporation)

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 5717106		19980210	
APPLICATION INFO.:	US 1996-724291		19960916	(8)

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Ramsuer, Robert W. LEGAL REPRESENTATIVE: Cole, Harold E.

NUMBER OF CLAIMS: 15 EXEMPLARY CLAIM: 447 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A yellow dye having the formula: ##STR1## wherein: R.sup.1, R.sup.2, R.sup.3 and R.sup.4 each independently represents hydrogen, halogen, or an alkoxy group of from 1 to about 6 carbon atoms; and

Z.sup.1 and Z.sup.2 each independently represents cyano, esterified carboxy, amide, a substituted or unsubstituted benzoxazole, or alkylsulfonyl; or may be taken together to form a pyrazolone, barbituric acid or Meldrum's acid residue.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 120 OF 122 USPATFULL on STN ACCESSION NUMBER: 85:53689 USPATFULL <<LOGINID::20110427>>

Article identification material and method and TITLE:

apparatus for using it INVENTOR(S): Acitelli, Mario A., Charlotte, NC, United States

Tynan, Richard F., Charlotte, NC, United States Wayson, Alan R., Concord, NC, United States

PATENT ASSIGNEE(S): International Business Machines Corporation, Armonk,

NY, United States (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: US 4540595 US 1982-433311 19850910 19821007 (6) APPLICATION INFO.:

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1982-344667, filed

on 1 Feb 1982, now abandoned

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Lusignan, Michael R. LEGAL REPRESENTATIVE: Coffman, E. Ronald

NUMBER OF CLAIMS: 10

EXEMPLARY CLAIM: NUMBER OF DRAWINGS: 5 Drawing Figure(s); 3 Drawing Page(s)

LINE COUNT: 329

CAS INDEXING IS AVAILABLE FOR THIS PATENT. AB

An ink that fluoresces in the near infrared is used to mark documents such as bank checks for automatic identification. Markings with this ink are reliably detectable, even in the presence of other markings commonly found on such documents. The preferred fluorescent material of our invention is a phenoxazine dye 3,7-BIS(diethylamino) phenoxazonium nitrate.

### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 121 OF 122 USPATFULL on STN

ACCESSION NUMBER: 73:6044 USPATFULL <<LOGINID::20110427>>

TITLE: INSPECTION PENETRANT DEVELOPMENT PROCESS EMPLOYING

FUSTBLE WAXES

INVENTOR(S): Alburger, James R., 5007 Hillard Avenue, La Canada, CA, United States 91011

NUMBER KIND DATE PATENT INFORMATION: APPLICATION INFO.: US 3715227 19730206 US 1971-127181 19710323 (5)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1969-799701, filed on 17 Feb 1969, now patented, Pat. No. US 3607333

DOCUMENT TYPE: Utility

FILE SEGMENT: Granted
PRIMARY EXAMINER: Leavitt, Alfred L.

ASSISTANT EXAMINER: Esposito, M. F. NUMBER OF CLAIMS:

NUMBER OF DRAWINGS: 2 Drawing Figure(s); 1 Drawing Page(s) LINE COUNT: 703

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

An improved process of inspection penetrant development employing a high-sensitivity, high-resolving power inspection penetrant developer in which the active developing ingredient is a waxy substance which is a solid or near-solid at room temperature but which becomes fluid at slightly elevated temperatures. The waxy developer material mat be dissolved in a suitable carrier liquid, such as water or other inert volatile solvent, and is deposited on test parts by dipping, brushing or spraying, and allowing the carrier liquid to evaporate. The development process includes the step of applying heat to the test parts, during oven drying or by hearing subsequent to air-drying, whereby the waxy developer layer becomes a fluid, and carries into solution any dyed penetrant entrapments present in the surface defects. When the test parts cool to room temperature, the fluid waxy layer, which now contains developed defect indications, solidifies and prevents excessive bleeding and migration of the indications.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 122 OF 122 USPATFULL on STN

ACCESSION NUMBER: 71:31997 USPATFULL <<LOGINID::20110427>>

TITLE: DEVELOPERS FOR INSPECTION PENETRANTS EMPLOYING FUSIBLE

WAXES INVENTOR(S): Alburger, James R., 5007 Hillard Ave., La Canada, CA,

United States 91011

NUMBER KIND DATE PATENT INFORMATION: US 3607333 19710921 APPLICATION INFO.: US 1969-799701 19690217 (4) DOCUMENT TYPE: Utility AFFICATION TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Liebman, Morris
ASSISTANT EXAMINER: Michl, P. R. NUMBER OF CLAIMS: 2 Drawing Figure(s); 1 Drawing Page(s) 628

NUMBER OF DRAWINGS: LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A high-sensitivity, high-resolving power inspection penetrant developer in which the active developing ingredient is a waxy substance which is a solid or near-solid at room temperature, but which becomes fluid at slightly elevated temperatures. The waxy developer material may be dissolved in a suitable carrier liquid such as water or other inert volatile solvent, and is deposited on test parts by dipping, brushing or spraying, and allowing the carrier liquid to evaporate. When heat is

applied to the test parts, during oven drying or by heating subsequent to air-drying, the waxy developer layer becomes a fluid, and a "liquid-film dilution-expansion" type development of penetrant entrapments in surface defects then takes place. When the test parts cool to room temperature, the fluid waxy layer, which now contains developed defect indications, solidifies and prevents excessive bleeding and migration of the indications.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

- => s (fluorescen? (3a) dye) (P) (masking (3a) background (5a) dye)
  PROXIMITY OPERATOR LEVEL NOT CONSISTENT MITH
  FIELD COOB 'AND' OPERATOR ASSUMED 'DYE) (P) '
  PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
  FIELD CODE 'AND' OPERATOR ASSUMED 'DYE) (P) '
  16 (FLUORESCEN? (3A) DYE) (P) (MASKING (3A) BACKGROUND (5A) DYE)
  => d 13 1-6 ti
- L3 ANSMER 1 OF 6 CAPLUS COPYRIGHT 2011 ACS on STN TI Masking background fluorescence and luminescence in optical analysis of biomedical assays
- L3 ANSWER 2 OF 6 USPATFULL on STN
- TI MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN THE OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS
- L3 ANSWER 3 OF 6 USPATFULL on STN
- TI Masking of the background fluorescence and luminescence in the optical analysis of biomedical assays
- L3 ANSWER 4 OF 6 USPATFULL on SIN
- TI MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS
- L3 ANSWER 5 OF 6 USPATFULL on STN
- TI Masking background fluorescence and luminescence in optical analysis of biomedical assays
- L3 ANSWER 6 OF 6 USPATFULL on STN
- TI Masking of the background fluorescence and luminescence in the optical analysis of biomedical assays
- => s (fluorescen? (3a) dye) (P) (second (3a) dye) (P) (reduc? (3a) background)
  PROXIMITY DEBRATOR LEVEL NOT CONSISTENT WITH
  FIELD CODE 'AND' OPERATOR ASSUMED 'DYE) (P) '
  PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
  FIELD CODE 'AND' OPERATOR ASSUMED 'DYE) (P) '
  PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
  FIELD CODE 'AND' OPERATOR ASSUMED 'DYE) (P) '
  PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
  FIELD CODE 'AND' OPERATOR ASSUMED 'DYE) (P)
  FROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
  FIELD CODE 'AND' OPERATOR ASSUMED 'DYE) (P)
  L4
  4 (FLUORESCEN? (3A) DYE) (P) (SECOND (3A) DYE) (P) (REDUC? (3A)
  BACKGROUND)
- => d 14 ti
- L4 ANSWER 1 OF 4 USPATFULL on STN
- TI Nucleic acid probes and methods to detect and/or quantify nucleic acid

## => d 14 1-4 ti

- ANSWER 1 OF 4 USPATFULL on STN L4
- Nucleic acid probes and methods to detect and/or quantify nucleic acid analytes
- ANSWER 2 OF 4 USPATFULL on STN L4
- TI Nucleic acid probes and methods to detect and/or quantify nucleic acid analytes
- ANSWER 3 OF 4 USPATFULL on STN T. 4
- TT Fluorescence digital imaging microscopy system
- ANSWER 4 OF 4 USPATFULL on STN T. 4
- TT Fluorescence digital imaging microscopy system

## => d 14 1-14 ibib abs

L4 ANSWER 1 OF 4 USPATFULL on STN

2005:268044 USPATFULL <<LOGINID::20110427>> ACCESSION NUMBER:

TITLE: Nucleic acid probes and methods to detect and/or quantify nucleic acid analytes

INVENTOR(S): Davies, Martin, Kent, UNITED KINGDOM

Bruce, Ian, East Sussex, UNITED KINGDOM

Wolter, Andreas, Esmarchstrasse, GERMANY, FEDERAL

(11)

REPUBLIC OF PATENT ASSIGNEE(S): PROLIGO, LLC, Boulder, CO, UNITED STATES (non-U.S.

corporation)

NUMBER KIND DATE PATENT INFORMATION: US 20050233360 A1 20051020 US 2005-83210 A1 20050316

RELATED APPLN. INFO.: Division of Ser. No. US 2002-278047, filed on 21 Oct 2002, GRANTED, Pat. No. US 6902900

NUMBER

PRIORITY INFORMATION: US 2001-336432P 20011019 (60)

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: SWANSON & BRATSCHUN L.L.C., 1745 SHEA CENTER DRIVE, SUITE 330, HIGHLANDS RANCH, CO, 80129, US

NUMBER OF CLAIMS: 20 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 21 Drawing Page(s)

LINE COUNT: 3448

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention comprises novel methods and strategies to detect and/or quantify nucleic acid analytes. The methods involve nucleic acid probes with covalently conjugated dyes, which are attached either at adjacent nucleotides or at the same nucleotide of the probe and novel linker molecules to attach the dyes to the probes. The nucleic acid probes generate a fluorescent signal upon hybridization to complementary nucleic acids based on the interaction of one of the attached dyes, which is either an intercalator or a DNA groove binder, with the formed double stranded DNA. The methods can be applied to a variety of

applications including homogeneous assays, real-time PCR monitoring, transcription assays, expression analysis on nucleic acid microarrays and other microarray applications such as genotyping (SNP analysis). The methods further include pH-sensitive nucleic acid probes that provide switchable fluorescence signals that are triggered by a change in the pH of the medium.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L4 ANSWER 2 OF 4 USPATFULL on STN

ACCESSION NUMBER: 2003:207233 USPATFULL <<LOGINID::20110427>>

TITLE: Nucleic acid probes and methods to detect and/or

quantify nucleic acid analytes
INVENTOR(S): Davies, Martin, Kent, UNITED KINGDOM

INVENTOR(S): Davies, Martin, Kent, UNITED KINGDOM
Bruce, Ian, East Sussex, UNITED KINGDOM

Wolter, Andreas, Hamburg, GERMANY, FEDERAL REPUBLIC OF

PATENT ASSIGNEE(S): PROLIGO, LLC, Boulder, CO, UNITED STATES, 80301

(non-U.S. corporation)

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 20030143591	A1	20030731	
	US 6902900	B2	20050607	
APPLICATION INFO.:	US 2002-278047	A1	20021021	(10

NUMBER DATE

20011019 (60)

PRIORITY INFORMATION: US 2001-336432P DOCUMENT TYPE: Utility

FILE SEGMENT: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: SWANSON & BRATSCHUN L.L.C., 1745 SHEA CENTER DRIVE,

SUITE 330, HIGHLANDS RANCH, CO, 80129

NUMBER OF CLAIMS: 60

EXEMPLARY CLAIM: 1
NUMBER OF DRAWINGS: 21 Drawing Page(s)

LINE COUNT: 3575

AB

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention comprises novel methods and strategies to detect and/or quantify nucleic acid analytes. The methods involve nucleic acid probes with covalently conjugated dyes, which are attached either at adjacent nucleotides or at the same nucleotide of the probe and novel linker molecules to attach the dyes to the probes. The nucleic acid probes generate a fluorescent signal upon hybridization to complementary nucleic acids based on the interaction of one of the attached dyes, which is either an intercalator or a DNA groove binder, with the formed double stranded DNA. The methods can be applied to a variety of applications including homogeneous assays, real-time PCR monitoring, transcription assays, expression analysis on nucleic acid microarrays and other microarray applications such as genotyping (SNP analysis). The methods further include pH-sensitive nucleic acid probes that provide switchable fluorescence signals that are triggered by a change in the pH of the medium.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L4 ANSWER 3 OF 4 USPATFULL on STN

ACCESSION NUMBER: 2002:336474 USPATFULL <<LOGINID::20110427>>

TITLE: Fluorescence digital imaging microscopy system

INVENTOR(S): Reynolds, C. Patrick, Sherman Oaks, CA, UNITED STATES Frgala, Tomas, Brno, CZECH REPUBLIC

PATENT ASSIGNEE(S): CHILDRENS HOSPITAL LOS ANGELES (U.S. corporation)

NUMBER KIND DATE US 20020191824 A1 20021219 PATENT INFORMATION: US 6665430 B2 20031216 A1 20020813 (10) US 2002-217721 APPLICATION INFO.: RELATED APPLN. INFO.: Division of Ser. No. US 1998-66134, filed on 24 Apr 1998, GRANTED, Pat. No. US 6459805 Continuation-in-part of Ser. No. US 1996-622110, filed on 26 Mar 1996, ABANDONED DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION LEGAL REPRESENTATIVE: HOGAN & HARTSON L.L.P., 500 S. GRAND AVENUE, SUITE 1900, LOS ANGELES, CA, 90071-2611 NUMBER OF CLAIMS: 14 EXEMPLARY CLAIM: NUMBER OF DRAWINGS: 13 Drawing Page(s) LINE COUNT: AB A method of preparing cell samples for viable cell number quantification with a fluorescence digital imaging microscopy system employing digital thresholding technique. The cell sample is stained with a first, fluorescent dve and treated with a second dve that is able to guench the fluorescence of the first dve. The fluorescent dve accumulates in viable cells only and is used to stain the viable cells. The second dye is excluded from viable cells but enters non-viable cells, thereby quenching the background fluorescence in non-viable cells and the medium. Two examples of dye combinations are described: fluorescein diacetate used as the fluorescent dye with eosin Y as the quenching dye; and calcein-AM used as the fluorescent dye with trypan blue as the quenching dye. By reducing the background fluorescence, the dynamic range and accuracy of viable cell number measurements are enhanced. In low viability cultures treated with

L4 ANSWER 4 OF 4 USPATFULL on STN

ACCESSION NUMBER: 2002:255074 USPATFULL <LOCINID::20110427>>
ITILE: Fluorescence digital imaging microscopy system
INVENTOR(S): Reynolds, C. Patrick, Sherman Oaks, CA, United States
Frgala, Tomas, Brno, CZECH REPUBLIC
Children Hospital Los Angeles, Los Angeles, CA, United

fluorescein diacetate, background fluorescence completely masked viable cells, but digital thresholding and eosin treatment dramatically

States (U.S. corporation)

reduced background fluorescence, producing a linear response over 4 logs of viable cell density.

	NUMBER	KIND	DATE		
PATENT INFORMATION:	US 6459805	B1	20021001		
APPLICATION INFO.:	US 1998-66134		19980424	(9)	
RELATED APPLN. INFO.:	Continuation-in-	part of	Ser. No.	US 1996-622110,	filed
	on 26 Mar 1996,	now abai	ndoned		
DOCUMENT TYPE:	Utility				
FILE SEGMENT:	GRANTED				
PRIMARY EXAMINER:	Patel, Jayanti K				
LEGAL REPRESENTATIVE:	Hogan & Hartson,	LLP			

NUMBER OF CLAIMS: 13 EXEMPLARY CLAIM: 1

NUMBER OF DRAWINGS: 19 Drawing Figure(s); 13 Drawing Page(s)

```
LINE COUNT: 70
```

=> d 16 1-7 ibib ti

AB

A method of preparing cell samples for viable cell number quantification with a fluorescence digital imaging microscopy system employing digital thresholding technique. The cell sample is stained with a first, fluorescent dve and treated with a second dye that is able to quench the fluorescence of the first dye. The fluorescent dye accumulates in viable cells only and is used to stain the viable cells. The second dve is excluded from viable cells but enters non-viable cells, thereby quenching the background fluorescence in non-viable cells and the medium. Two examples of dve combinations are described: fluorescein diacetate used as the fluorescent dye with eosin Y as the quenching dye; and calcein-AM used as the fluorescent dye with trypan blue as the quenching dye. By reducing the background fluorescence, the dynamic range and accuracy of viable cell number measurements are enhanced. In low viability cultures treated with fluorescein diacetate, background fluorescence completely masked viable cells, but digital thresholding and eosin treatment dramatically reduced background fluorescence, producing a linear response over 4 logs of viable cell density.

```
=> e reynolds c patrick/au
           1 REYNOLDS C PARTRIK/AU
E1
E2
                  REYNOLDS C PATRIC/AU
E3
           349 --> REYNOLDS C PATRICK/AU
E4
           1 REYNOLDS C PATRIK/AU
                 REYNOLDS C R/AU
         136
246
E5
                 REYNOLDS C S/AU
E6
E7
          29
                 REYNOLDS C T/AU
E8
                 REYNOLDS C T 3RD/AU
          1 REYNOLDS C T JR/AU
7 REYNOLDS C V/AU
275 REYNOLDS C W/AU
E9
E10
E11
E12
           3
                  REYNOLDS CALVIN E/AU
=> s e3 and (dye)
           12 "REYNOLDS C PATRICK"/AU AND (DYE)
=> dup remm 15
ENTER REMOVE, IDENTIFY, ONLY, OR (?):dup rem 15
'DUP REM L41' IS NOT VALID HERE
Enter "REMOVE" to identify and remove duplicate answers.
Enter "IDENTIFY" to identify duplicate answers in the answer set.
Enter "ONLY" to identify and create an answer set containing only
duplicate records.
ENTER REMOVE, IDENTIFY, ONLY, OR (?):?
Enter "REMOVE" to identify and remove duplicate answers.
Enter "IDENTIFY" to identify duplicate answers in the answer set.
Enter "ONLY" to identify and create an answer set containing only
duplicate records.
ENTER REMOVE, IDENTIFY, ONLY, OR (?):end
=> dup rem 15
PROCESSING COMPLETED FOR L5
              7 DUP REM L5 (5 DUPLICATES REMOVED)
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L6 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN DUPLICATE 1

ACCESSION NUMBER: 2007:289476 CAPLUS <<LOGINID::20110427>> 146:394606

DOCUMENT NUMBER:

TITLE: A fluorescence microplate cytotoxicity assay with a 4-log dynamic range that identifies synergistic drug

combinations

Frgala, Tomas; Kalous, Ondrej; Proffitt, Robert T.; AUTHOR(S):

Reynolds, C. Patrick

CORPORATE SOURCE: Developmental Therapeutics Program, USC-CHLA Institute for Pediatric Clinical Research, Childrens Hospital of

Los Angeles and Division Hematology-Oncology,

Department of Pediatrics, The University of Southern

California Keck School of Medicine, Los Angeles, CA, 90027, USA

SOURCE: Molecular Cancer Therapeutics (2007), 6(3), 886-897

CODEN: MCTOCF; ISSN: 1535-7163

PHRLISHER. American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English

A fluorescence microplate cytotoxicity assay with a 4-log dynamic range

that identifies synergistic drug combinations OS.CITING REF COUNT: THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD

(7 CITINGS)

REFERENCE COUNT: 60 THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN DUPLICATE 2 ACCESSION NUMBER: 2007:792406 CAPLUS <<LOGINID::20110427>>

DOCUMENT NUMBER: 147:132289

TITLE: Assessing combinations of cytotoxic agents using

leukemia cell lines

AUTHOR(S): Reynolds, C. Patrick; Kang, Min H.; Keshelava, Nino; Maurer, Barry J.

CORPORATE SOURCE: Developmental Therapeutics Program, USC-CHLA Institute

for Pediatric Clinical Research and Division of Hematology-Oncology, Department of Pediatrics, Keck School of Medicine, University of Southern California

and Childrens Hospital Los Angeles, USA SOURCE: Current Drug Targets (2007), 8(6), 765-771

CODEN: CDTUAU; ISSN: 1389-4501

PUBLISHER: Bentham Science Publishers Ltd.

DOCUMENT TYPE: Journal: General Review

LANGHAGE · English

TI Assessing combinations of cytotoxic agents using leukemia cell lines OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD

(2 CITINGS)

REFERENCE COUNT: THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS 41 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2011 ACS on STN DUPLICATE 3 ACCESSION NUMBER: 2005:419086 CAPLUS <<LOGINID::20110427>>

DOCUMENT NUMBER: 144:120778

TITLE: DIMSCAN a microcomputer fluorescence-based

cytotoxicity assay for preclinical testing of

combination chemotherapy Keshelava, Nino; Frgala, Tomas; Krejsa, Jiri; Kalous, AUTHOR(S): Ondrej; Reynolds, C. Patrick

CORPORATE SOURCE: USC-CHLA Institute for Pediatric Clinical Research,

University of Southern California and Childrens Hospital Los Angeles, Los Angeles, CA, USA

SOURCE . Methods in Molecular Medicine (2005), 110 (Chemosensitivity, Volume 1), 139-153

CODEN: MMMEEN

PUBLISHER: Humana Press Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

TI DIMSCAN a microcomputer fluorescence-based cytotoxicity assay for

preclinical testing of combination chemotherapy

OS.CITING REF COUNT: 10 THERE ARE 10 CAPLUS RECORDS THAT CITE THIS

RECORD (10 CITINGS)

37 REFERENCE COUNT: THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 7 USPATFULL on STN

ACCESSION NUMBER: 2002:336474 USPATFULL <<LOGINID::20110427>> TITLE: Fluorescence digital imaging microscopy system

INVENTOR(S): Reynolds, C. Patrick, Sherman Oaks, CA,

UNITED STATES

Frgala, Tomas, Brno, CZECH REPUBLIC

PATENT ASSIGNEE(S): CHILDRENS HOSPITAL LOS ANGELES (U.S. corporation)

NUMBER KIND DATE US 20020191824 A1 20021219 PATENT INFORMATION: B2 20031216 US 6665430 US 2002-217721 APPLICATION INFO.: A1 20020813 (10)

RELATED APPLN. INFO.: Division of Ser. No. US 1998-66134, filed on 24 Apr

1998, GRANTED, Pat. No. US 6459805 Continuation-in-part of Ser. No. US 1996-622110, filed on 26 Mar 1996,

ABANDONED

Utility DOCUMENT TYPE: FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: HOGAN & HARTSON L.L.P., 500 S. GRAND AVENUE, SUITE

1900, LOS ANGELES, CA, 90071-2611

NUMBER OF CLAIMS: 14

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

13 Drawing Page(s) LINE COUNT:

Fluorescence digital imaging microscopy system

L6 ANSWER 5 OF 7 USPATFULL on STN

ACCESSION NUMBER: 2002:255074 USPATFULL <<LOGINID::20110427>> TITLE: Fluorescence digital imaging microscopy system

INVENTOR(S): Reynolds, C. Patrick, Sherman Oaks, CA,

United States Frgala, Tomas, Brno, CZECH REPUBLIC

Childrens Hospital Los Angeles, Los Angeles, CA, United PATENT ASSIGNEE(S):

States (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: US 6459805 B1 20021001 US 1998-66134 APPLICATION INFO.: 19980424 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1996-622110, filed

on 26 Mar 1996, now abandoned

DOCUMENT TYPE: Utility

FILE SEGMENT: GRANTED PRIMARY EXAMINER: Patel, Jayanti K.

LEGAL REPRESENTATIVE: Hogan & Hartson, LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 19 Drawing Figure(s); 13 Drawing Page(s) LINE COUNT: 705

TI Fluorescence digital imaging microscopy system

L6 ANSWER 6 OF 7 USPATFULL on STN

ACCESSION NUMBER: 2002:75229 USPATFULL <<LOGINID::20110427>> Treatment of hyperproliferative disorders TITLE: INVENTOR(S): Maurer, Barry J., Sunland, CA, United States

Reynolds, C. Patrick, Sherman Oaks, CA,

United States

PATENT ASSIGNEE(S): Childrens Hospital Los Angeles, Los Angeles, CA, United

States (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: US 6368831 B1 20020409 APPLICATION INFO.: US 1999-471944 19991223 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1999-342019, filed

on 28 Jun 1999

NUMBER DATE PRIORITY INFORMATION: US 1998-91138P 19980629 (60)

DOCUMENT TYPE: Utility

FILE SEGMENT: GRANTED PRIMARY EXAMINER: ASSISTANT EXAMINER: Low, Christopher S. F.

Robinson, Hope A.

LEGAL REPRESENTATIVE: Myers Bigel Sibley & Sajovec 30

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

21 Drawing Figure(s); 16 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 1466

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Treatment of hyperproliferative disorders

L6 ANSWER 7 OF 7 USPATFULL on STN

ACCESSION NUMBER: 2002:45484 USPATFULL <<LOGINID::20110427>> TITLE: Treatment of hyperproliferative disorders INVENTOR(S): Maurer, Barry J., Pasadena, CA, United States Cabot, Myles, Santa Monica, CA, United States

Reynolds, C. Patrick, Sherman Oaks, CA, United States

PATENT ASSIGNEE(S): Childrens Hospital Los Angeles, Los Angeles, CA, United

States (U.S. corporation)

John Wayne Cancer Institute, Santa Monica, CA, United

States (U.S. corporation)

NUMBER KIND DATE US 6352844 B1 20020305 US 1999-342019 19990628 PATENT INFORMATION: APPLICATION INFO.: 19990628 (9)

NUMBER US 1998-91138P 19980629 (60) PRIORITY INFORMATION:

DOCUMENT TYPE: Utility FILE SEGMENT: GRANTED

PRIMARY EXAMINER: GARRIED Carlson, Karen Cochrane ASSISTANT EXAMINER: Robinson, Hope A.

LEGAL REPRESENTATIVE: Myers Bigel Sibley & Sajovec, P.A. NUMBER OF CLAIMS: 19

EXEMPLARY CLAIM:

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NUMBER OF DRAWINGS:
                     21 Drawing Figure(s); 15 Drawing Page(s)
LINE COUNT:
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Treatment of hyperproliferative disorders
=> e frgala tomas/au
           13
                 FRGALA JIRI/AU
E2
           21
                  FRGALA T/AU
E3
           23 --> FRGALA TOMAS/AU
E4
           15 FRGALA Z/AU
E5
           1
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E6
          18
                 FRGALOVA K/AU
          12
E7
                 FRGALOVA KARLA/AU
E8
           2
                 FRGEMAN NILS J/AU
E9
            1
                 FRGEMAN O/AU
E10
            3
                 FRGEMAND MERETE/AU
                FRGESTAD ELLEN M/AU
FRGESTAD ELLEN MOSLETH/AU
E11
            1
E12
            1
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L7
             7 "FRGALA TOMAS"/AU AND DYE
=> dup rem 17
PROCESSING COMPLETED FOR L7
             4 DUP REM L7 (3 DUPLICATES REMOVED)
=> d 18 1-4
    ANSWER 1 OF 4 CAPLUS COPYRIGHT 2011 ACS on STN DUPLICATE 1
T.R
    2007:289476 CAPLUS <<LOGINID::20110427>>
AN
DN
    146:394606
    A fluorescence microplate cytotoxicity assay with a 4-log dynamic range
    that identifies synergistic drug combinations
    Frgala, Tomas; Kalous, Ondrej; Proffitt, Robert T.; Reynolds, C.
ΑU
    Patrick
    Developmental Therapeutics Program, USC-CHLA Institute for Pediatric
    Clinical Research, Childrens Hospital of Los Angeles and Division
     Hematology-Oncology, Department of Pediatrics, The University of Southern
    California Keck School of Medicine, Los Angeles, CA, 90027, USA
    Molecular Cancer Therapeutics (2007), 6(3), 886-897
    CODEN: MCTOCF; ISSN: 1535-7163
PB
   American Association for Cancer Research
DT
    Journal
LA
   English
OSC.G 7
             THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD (7 CITINGS)
RE.CNT 60
             THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
L8
    ANSWER 2 OF 4 CAPLUS COPYRIGHT 2011 ACS on STN DUPLICATE 2
AN
    2005:419086 CAPLUS <<LOGINID::20110427>>
DN
     144:120778
     DIMSCAN a microcomputer fluorescence-based cytotoxicity assay for
     preclinical testing of combination chemotherapy
    Keshelava, Nino; Frgala, Tomas; Krejsa, Jiri; Kalous, Ondrej;
    Reynolds, C. Patrick
    USC-CHLA Institute for Pediatric Clinical Research, University of Southern
    California and Childrens Hospital Los Angeles, Los Angeles, CA, USA
SO
    Methods in Molecular Medicine (2005), 110 (Chemosensitivity, Volume 1),
     139-153
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CODEN: MMMEFN

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PB
    Humana Press Inc.
DT
    Journal
LA
    English
      10
             THERE ARE 10 CAPLUS RECORDS THAT CITE THIS RECORD (10 CITINGS)
OSC.G
RE.CNT 37
             THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD
             ALL CITATIONS AVAILABLE IN THE RE FORMAT
L8
    ANSWER 3 OF 4 USPATFULL on STN
AN
       2002:336474 USPATFULL <<LOGINID::20110427>>
       Fluorescence digital imaging microscopy system
TI
IN
       Reynolds, C. Patrick, Sherman Oaks, CA, UNITED STATES
         Frgala, Tomas, Brno, CZECH REPUBLIC
PA
       CHILDRENS HOSPITAL LOS ANGELES (U.S. corporation)
PΙ
       US 20020191824
                          A1 20021219
      US 6665430
                           B2 20031216
AΤ
      US 2002-217721
                          A1 20020813 (10)
RLT
      Division of Ser. No. US 1998-66134, filed on 24 Apr 1998, GRANTED, Pat.
      No. US 6459805 Continuation-in-part of Ser. No. US 1996-622110, filed on
       26 Mar 1996, ABANDONED
      Utility
      APPLICATION
LN.CNT 712
       INCLM: 382/128.000
INCL
NCL
      NCLM: 382/128.000
      NCLS: 435/006.000; 435/029.000; 435/040.500
TPC:
       TPCT
             G06K0009-00 [ICM, 7]
       IPCI-2 G06K0009-00 [ICM, 7]
       IPCR G01N0021-64 [I,C*]; G01N0021-64 [I,A]
    ANSWER 4 OF 4 USPATFULL on STN
T.R
AN
       2002:255074 USPATFULL <<LOGINID::20110427>>
       Fluorescence digital imaging microscopy system
       Reynolds, C. Patrick, Sherman Oaks, CA, United States
        Frgala, Tomas, Brno, CZECH REPUBLIC
PA
      Childrens Hospital Los Angeles, Los Angeles, CA, United States (U.S.
       corporation)
PΙ
      US 6459805
                           B1 20021001
ΑI
      US 1998-66134
                              19980424 (9)
RLT
      Continuation-in-part of Ser. No. US 1996-622110, filed on 26 Mar 1996,
      now abandoned
DT
      Utility
FS
      GRANTED
LN.CNT 705
INCL
       INCLM: 382/128.000
       INCLS: 435/029.000; 435/040.500; 436/172.000
NCL.
      NCLM: 382/128.000
      NCLS: 435/029.000; 435/040.500; 436/172.000
TPC
       IPCI
             G06K0009-00 [ICM, 7]
             G01N0021-64 [I,C*]; G01N0021-64 [I,A]
       382/100; 382/128; 382/129-134; 382/312; 435/6; 435/29; 435/34; 435/40.5;
       435/325; 435/968; 435/177; 430/138-139; 436/172-173; 356/39-42; 514/440;
       514/629; 514/634; 549/33
=> s ( fluorescent (3a) dye) (p) (brilliant (3a) black)
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
FIELD CODE - 'AND' OPERATOR ASSUMED 'DYE) (P) '
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
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FIELD CODE - 'AND' OPERATOR ASSUMED 'DYE) (P) '

=> dup rem 19

PROCESSING COMPLETED FOR L9

L10 29 DUP REM L9 (0 DUPLICATES REMOVED)

=> d 110 1-29 t.i

L10 ANSWER 1 OF 29 USPATFULL on STN

TI METHOD FOR MEASURING MITOCHONDRIAL MEMBRANE POTENTIAL IN VERTEBRATE CELLS

L10 ANSWER 2 OF 29 USPATFULL on STN

TI POLYMERIZED CONJUGATES FOR BIOLOGICAL APPLICATIONS

L10 ANSWER 3 OF 29 USPATFULL on STN

TI MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN THE OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS

L10 ANSWER 4 OF 29 USPATFULL on STN

TI INFRARED TRANSMISSIVE THERMOPLASTIC COMPOSITION, AND ARTICLES FORMED THEREFROM

L10 ANSWER 5 OF 29 USPATFULL on STN

TI Methods for Production of Proteins

L10 ANSWER 6 OF 29 USPATFULL on STN

'I Methods for Production of Proteins

L10 ANSWER 7 OF 29 USPATFULL on STN

TI Methods for production of proteins

L10 ANSWER 8 OF 29 USPATFULL on STN

TI Methods of production of proteins

L10 ANSWER 9 OF 29 USPATFULL on STN

II Water-soluble conjugates for electrochemical detection

L10 ANSWER 10 OF 29 USPATFULL on STN

T Water-soluble conjugates for electrochemical detection

L10 ANSWER 11 OF 29 USPATFULL on STN

TI Water-soluble conjugates and methods of preparation

L10 ANSWER 12 OF 29 USPATFULL on STN

TI Use of indole-3-acetic acids in the treatment of asthma, copd and other diseases

L10 ANSWER 13 OF 29 USPATFULL on STN

TI Use of indole-3-acetic acids in the treatment of asthma, copd and other diseases

L10 ANSWER 14 OF 29 USPATFULL on STN

II Coating compositions and processes

L10 ANSWER 15 OF 29 USPATFULL on STN

TI Methods for production of proteins

L10 ANSWER 16 OF 29 USPATFULL on STN

TI Lipoprotein fingerprinting methods using metal ion chelate salts

- L10 ANSWER 17 OF 29 USPATFULL on STN
- TI Masking of the background fluorescence and luminescence in the optical analysis of biomedical assays
- L10 ANSWER 18 OF 29 USPATFULL on STN
- II Lipoprotein fingerprinting method
- L10 ANSWER 19 OF 29 USPATFULL on STN
- TI Method for preparing water-soluble cross-linked conjugates
- L10 ANSWER 20 OF 29 USPATFULL on STN
- TI METHODS FOR PRODUCTION OF PROTEIN
- L10 ANSWER 21 OF 29 USPATFULL on STN
- TI MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS
- L10 ANSWER 22 OF 29 USPATFULL on STN
- TI Masking background fluorescence and luminescence in optical analysis of biomedical assays
- L10 ANSWER 23 OF 29 USPATFULL on STN
- TI Masking of the background fluorescence and luminescence in the optical analysis of biomedical assays
- L10 ANSWER 24 OF 29 USPATFULL on STN
- TI Waterfast ink jet inks containing an emulsifiable polymer resin
- L10 ANSWER 25 OF 29 USPATFULL on STN
- TI Process for making multilayer coatings with a strippable topcoat
- L10 ANSWER 26 OF 29 CAPLUS COPYRIGHT 2011 ACS on STN
- TI Complexities of measuring antagonist potency at P2X7 receptor orthologs
- L10 ANSWER 27 OF 29 USPATFULL on STN
  - Colored particulates for ink jet inks
- L10 ANSWER 28 OF 29 CAPLUS COPYRIGHT 2011 ACS on STN
- II Non-toxic, water-soluble photocalorimetric reference compounds for UV and visible excitation
- L10 ANSWER 29 OF 29 CAPLUS COPYRIGHT 2011 ACS on STN
- TI Printing of knit fabrics
- => s (brilliant (3a) black) same (mask?) MISSING OPERATOR BLACK) SAME

The search profile that was entered contains terms or

nested terms that are not separated by a logical operator.

- => s (brilliant (3a) black) (p) (mask?)
  PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
- FIELD CODE 'AND' OPERATOR ASSUMED 'BLACK) (P) 'PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
- FIELD CODE 'AND' OPERATOR ASSUMED 'BLACK) (P) '
- L11 13 (BRILLIANT (3A) BLACK) (P) (MASK?)
- => dup rem 111
- PROCESSING COMPLETED FOR L11
- L12 13 DUP REM L11 (0 DUPLICATES REMOVED)

L12 ANSWER 1 OF 13 USPATFULL on STN

ACCESSION NUMBER: 2008:361999 USPATFULL <<LOGINID::20110427>>

TITLE: MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN THE OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS

INVENTOR(S): Krahn, Thomas, Hagen, GERMANY, FEDERAL REPUBLIC OF Paffhausen, Wolfgang, Leverkusen, GERMANY, FEDERAL

Paffhausen, Wolfgang, Leverkusen, GERMANY, FEDE

REPUBLIC OF

1997

Schade, Andreas, Essen, GERMANY, FEDERAL REPUBLIC OF Bechem, Martin, Wuppertal, GERMANY, FEDERAL REPUBLIC OF Schmidt, Deif, Wuppertal, GERMANY, FEDERAL REPUBLIC OF

PATENT ASSIGNEE(S): BAYER HEALTHCARE AG, Leverkusen, GERMANY, FEDERAL REPUBLIC OF (non-U.S. corporation)

NUMBER DATE
\_\_\_\_\_\_
DE 1996-19621312 19960528

PRIORITY INFORMATION: DOCUMENT TYPE:

DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: NORRIS, MCLAUGHLIN & MARCUS, PA, 875 THIRD AVENUE, 18TH FLOOR, NEW YORK, NY, 10022, US

NUMBER OF CLAIMS: 4

EXEMPLARY CLAIM: 1-6

NUMBER OF DRAWINGS: 10 Drawing Page(s)
LINE COUNT: 445

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3 containing the fluorescent dve 4. The sensitivity of analytical detection can be considerably improved if to the fluorescent dye 4 already present in addition a masking dye 9, which absorbs the excitation light 6 for the fluorescent dye 4 and/or its emission light 7, is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6 or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating layer 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously, these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor layer 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here if a masking dye 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission light or (in the case of luminescent

ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye in the solution 3 or optionally as an additional measure, a separating layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light can be applied to the cell or receptor layer 12 at the bottom 2.

#### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 2 OF 13 USPATFULL on STN

2005:158295 USPATFULL <<LOGINID::20110427>> ACCESSION NUMBER: TITLE: Fluorescent pH indicators for intracellular assays

INVENTOR(S): Diwu, Zhenjun, Sunnyvale, CA, UNITED STATES Twu, Jesse J., Cupertino, CA, UNITED STATES Yi, Guoliang, Sunnyvale, CA, UNITED STATES Lavis, Luke D., Sunnyvale, CA, UNITED STATES

Chen, Yen-Wen, San Francisco, CA, UNITED STATES Cassutt, Kelly J., Somerset, NJ, UNITED STATES

NUMBER KIND DATE US 20050136503 A1 20050623 US 7507395 B2 20090324 US 2004-958670 A1 20041004 (10) PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.: Continuation of Ser. No. US 2002-108656, filed on 27 Mar 2002, GRANTED, Pat. No. US 6800765

NUMBER DATE

US 2001-309800P 20010802 (60) PRIORITY INFORMATION:

DOCUMENT TYPE: Utility FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE: KOLISCH HARTWELL, P.C., 520 S.W. YAMHILL STREET, SUITE 200, PORTLAND, OR, 97204, US

23 NUMBER OF CLAIMS:

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 6 Drawing Page(s) LINE COUNT: 1160

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Systems, including compositions and methods, for measuring pH, particularly in cells, organelles, and other samples. The compositions

include pH-sensitive fluorescent and fluorogenic

2'.7'-dialkylfluorescein derivatives and associated nonfluorescent precursor compounds. The compositions may permit ratiometric measurement in the excitation spectrum and the emission spectrum. The methods include adding a precursor compound to a sample cell, incubating the sample cell to release the free indicator, illuminating the sample cell,

and detecting the fluorescence response of the free indicator.

CAS INDEXING IS AVAILABLE FOR THIS PATENT. L12 ANSWER 3 OF 13 USPATFULL on STN

ACCESSION NUMBER: 2003:133996 USPATFULL <<LOGINID::20110427>>

TITLE: Masking of the background fluorescence and luminescence

in the optical analysis of biomedical assays Krahn, Thomas, Hagen, GERMANY, FEDERAL REPUBLIC OF INVENTOR(S): Paffhausen, Wolfgang, Leverkusen, GERMANY, FEDERAL

REPUBLIC OF

Schade, Andreas, Essen, GERMANY, FEDERAL REPUBLIC OF Bechem, Martin, Wuppertal, GERMANY, FEDERAL REPUBLIC OF Schmidt, Delf, Wuppertal, GERMANY, FEDERAL REPUBLIC OF

NUMBER KIND DATE US 20030092081 A1 20030515 US 7138280 B2 20061121 US 2002-263607 A1 20021003 (10)

APPLICATION INFO.: Division of Ser. No. US 2001-966522, filed on 28 Sep RELATED APPLN. INFO.:

2001, PENDING

NUMBER DATE

PRIORITY INFORMATION: DE 1996-19621312 19960528 DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

PATENT INFORMATION:

LEGAL REPRESENTATIVE: KURT BRISCOE, NORRIS, MCLAUGHLIN & MARCUS, P.A., 220 EAST 42ND STREET, 30TH FLOOR, NEW YORK, NY, 10017

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 10 Drawing Page(s) LINE COUNT: 438

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3 containing the fluorescent dye 4. The sensitivity of analytical detection can be considerably improved if to the fluorescent dye 4 already present in addition a masking dye 9, which absorbs the excitation light 6 for the fluorescent dye 4 and/or its emission light 7, is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6 or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating layer 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously, these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor layer 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here if a masking dve 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission light or (in the case of luminescent ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye in the solution 3 or optionally as an additional measure, a separating layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light can be applied to the cell or receptor layer 12 at the bottom 2.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 4 OF 13 USPATFULL on STN

ACCESSION NUMBER: TITLE: INVENTOR(S):

2003:99563 USPATFULL <<LOGINID::20110427>> Fluorescent pH indicators for intracellular assays Diwu, Zhenjun, Sunnyvale, CA, UNITED STATES Twu, Jesse J., Cupertino, CA, UNITED STATES Yi, Guoliang, Sunnyvale, CA, UNITED STATES Lavis, Luke D., Sunnyvale, CA, UNITED STATES Chen, Yen-Wen, San Francisco, CA, UNITED STATES

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 20030068668	A1	20030410	
	US 6800765	B2	20041005	
APPLICATION INFO.:	US 2002-108656	A1	20020327	(10)
	NUMBER		DATE	

NUMBER PRIORITY INFORMATION: US 2001-309800P 20010802 (60)

DOCUMENT TYPE: Utility

FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: James R. Abney, Kolisch, Hartwell, Dickinson, McCormack

& Heuser, 200 Pacific Building, 520 S.W. Yamhill

Street, Portland, OR, 97204

NUMBER OF CLAIMS: 57

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 6 Drawing Page(s) LINE COUNT: 1298

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Systems, including compositions and methods, for measuring pH, particularly in cells, organelles, and other samples. The compositions include pH-sensitive fluorescent and fluorogenic

2'.7'-dialkylfluorescein derivatives and associated nonfluorescent precursor compounds. The compositions may permit ratiometric measurement in the excitation spectrum and the emission spectrum. The methods include adding a precursor compound to a sample cell, incubating the sample cell to release the free indicator, illuminating the sample cell,

and detecting the fluorescence response of the free indicator.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 5 OF 13 USPATFULL on STN

ACCESSION NUMBER: 2002:37557 USPATFULL <<LOGINID::20110427>>

TITLE: MASKING BACKGROUND FLUORESCENCE AND LUMINESCENCE IN OPTICAL ANALYSIS OF BIOMEDICAL ASSAYS

INVENTOR(S): KRAHN, THOMAS, HAGEN, GERMANY, FEDERAL REPUBLIC OF

PAFFHAUSEN, WOLFGANG, LEVERKUSEN, GERMANY, FEDERAL REPUBLIC OF

SCHADE, ANDREAS, ESSEN, GERMANY, FEDERAL REPUBLIC OF BECHEM, MARTIN, WUPPERTAL, GERMANY, FEDERAL REPUBLIC OF SCHMIDT, DELF, WUPPERTAL, GERMANY, FEDERAL REPUBLIC OF

	NUMBER	KIND	DATE	
PATENT INFORMATION:	US 20020022274	A1	20020221	
	US 6420183	B2	20020716	
APPLICATION INFO.:	US 1998-194099	A1	19981120	(9)
	WO 1997-EP2662		19970523	

		NUMBER	DATE
DDTADTTV	THEODMATION.	DE 1996-19621312	1996052

Utility DOCUMENT TYPE: APPLICATION

LEGAL REPRESENTATIVE: NORRIS McLAUGHLIN & MARCUS, P.A., 220 EAST 42nd STREET

30TH FLOOR, NEW YORK, NY, 10017

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 10 Drawing Page(s)

LINE COUNT: 462

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3 containing the fluorescent dye 4. The sensitivity of analytical detection can be considerably improved if to the fluorescent dye 4 already present in addition a masking dye 9, which absorbs the excitation light 6 for the fluorescent dve 4 and/or its emission light 7, is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6 or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating layer 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously, these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor layer 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here if a masking dve 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission light or (in the case of luminescent ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye in the solution 3 or optionally as an additional measure, a separating layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light can be applied to the cell or receptor layer 12 at

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 6 OF 13 USPATFULL on STN

the bottom 2.

ACCESSION NUMBER: 2002:27123 USPATFULL <<LOGINID::20110427>>

MINIOPO

TITLE: Masking background fluorescence and luminescence in

optical analysis of biomedical assays

INVENTOR(S): Krahn, Thomas, Hagen, GERMANY, FEDERAL REPUBLIC OF Paffhausen, Wolfgang, Leverkusen, GERMANY, FEDERAL

REPUBLIC OF

Schade, Andreas, Essen, GERMANY, FEDERAL REPUBLIC OF Bechem, Martin, Wuppertal, GERMANY, FEDERAL REPUBLIC OF Schmidt, Delf, Wuppertal, GERMANY, FEDERAL REPUBLIC OF DATE

	NUMBER	KIND	DATE
PATENT INFORMATION:	US 20020015969	A1	20020207
	US 7063952	B2	20060620
APPLICATION INFO.:	US 2001-966137	A1	20010928 (9)
RELATED APPLN. INFO.:	Division of Ser. 1998, PENDING	No. US	1998-194099, filed on 20 Nov

ECTNID

NUMBER	DATE

PRIORITY INFORMATION: DE 1996-19621312 19960528 DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION

LEGAL REPRESENTATIVE: Kurt G. Briscoe, Norris McLaughlin & Marcus, P.A., 30th Floor, 220 East 42nd Street, New York, NY, 10017

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: 1
NUMBER OF DRAWINGS: 10 Drawing Page(s)
LINE COUNT: 462

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3 containing the fluorescent dye 4. The sensitivity of analytical detection can be considerably improved if to the fluorescent dve 4 already present in addition a masking dye 9, which absorbs the excitation light 6 for the fluorescent dve 4 and/or its emission light 7, is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6 or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating layer 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously, these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor laver 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here if a masking dye 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission light or (in the case of luminescent ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye in the solution 3 or optionally as an additional measure, a separating layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light can be applied to the cell or receptor layer 12 at the bottom 2.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 7 OF 13 USPATFULL on STN

ACCESSION NUMBER: TITLE:

INVENTOR(S):

2002:16874 USPATFULL <<LOGINID::20110427>> Masking of the background fluorescence and luminescence in the optical analysis of biomedical assays Krahn, Thoams, Hagen, GERMANY, FEDERAL REPUBLIC OF Paffhausen, Wolfgang, Leverkusen, GERMANY, FEDERAL REPUBLIC OF Schade, Andreas, Essen, GERMANY, FEDERAL REPUBLIC OF Bechem, Martin, Wuppertal, GERMANY, FEDERAL REPUBLIC OF

Schmidt, Delf, Wuppertal, GERMANY, FEDERAL REPUBLIC OF NUMBER KIND DATE

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

US 20020009754 A1 20020124 US 2001-966522 A1 20010928 (9) Continuation of Ser. No. US 1998-194099, filed on 20 Nov 1998, PENDING

NUMBER DATE DE 1996-19621312 19960528 Utility

PRIORITY INFORMATION: DOCUMENT TYPE: FILE SEGMENT:

APPLICATION LEGAL REPRESENTATIVE: Kurt G. Briscoe, Norris McLaughlin & Marcus, P.A., 30th Floor, 220 East 42nd Street, New York, NY, 10017

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

10 Drawing Page(s) NUMBER OF DRAWINGS: 462

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

In a process for the quantitative optical analysis of fluorescently labelled biological cells 5, a cell layer on a transparent support at the bottom 2 of a reaction vessel 1 is in contact with a solution 3 containing the fluorescent dve 4. The sensitivity of analytical detection can be considerably improved if to the fluorescent dve 4 already present in addition a masking dye 9, which absorbs the excitation light 6 for the fluorescent dye 4 and/or its emission light 7, is added to the solution 3 and/or if a separating layer 10 permeable to the solution and absorbing and/or reflecting the excitation light 6 or the emission light 7 is applied to the cell layer at the bottom 2. This process can also be used for improving the sensitivity in the quantitative optical analysis of a luminescent biological cell layer. The separating layer 10 must in this case be composed such that it has a high power of reflection for the luminescent light 11. Analogously, these process principles can also be used in receptor studies for the masking of the interfering background radiation in the quantitative optical analysis of fluorescently or luminescently labelled reaction components. In this case, a receptor laver 12 at the bottom 2 of a reaction vessel 1 is in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand 13 is dissolved. The sensitivity and accuracy of the analytical detection can be considerably improved here if a masking dye 9 which absorbs the excitation light 6 for the fluorescent dye and/or its emission light or (in the case of luminescent ligands) the luminescent light is added to the supernatant 3. Instead of the masking dye in the solution 3 or optionally as an additional measure, a separating layer 10 permeable to the solution 3 and absorbing and/or reflecting the excitation light 6 and/or the emission light or the luminescent light can be applied to the cell or receptor layer 12 at

# the bottom 2. (FIGS. 2 and 10) CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 8 OF 13 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1997:805887 CAPLUS <<LOGINID::20110427>>

DOCUMENT NUMBER: 128:59162

ORIGINAL REFERENCE NO.: 128:11503a,11506a

TITLE: Masking background fluorescence and luminescence in

optical analysis of biomedical assays

INVENTOR(S): Krahn, Thomas; Paffhausen, Wolfgang; Schade, Andreas; Bechem, Martin; Schmidt, Delf

Bayer Aktiengesellschaft, Germany; Krahn, Thomas; PATENT ASSIGNEE(S):

Paffhausen, Wolfgang; Schade, Andreas; Bechem, Martin;

Schmidt, Delf

PCT Int. Appl., 30 pp. SOURCE .

CODEN: PIXXD2

DOCUMENT TYPE:

Patent German

LANGUAGE: FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	ENT	NO.			KIN	D	DATE		APPL	ICAT.	TON .	NO.		D	ATE		
						-											
WO	9745	739			A1		1997	1204	WO 1	997-1	EP26	62		1	9970	523	
	W:	CA,	JP,	US													
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FI,	FR, GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE
DE	1962	1312			A1		1997	1204	DE 1	996-	1962	1312		11	9960	528	

	2256629		A1	19971204	CA	1997-2256629		19970523
CA	2256629		C	20030722				
EP	906572		A1	19990407	EP	1997-927032		19970523
EP	906572		В1	20020403				
	R: AT, BE,	CH,	DE,	DK, ES, FR,	GB, I	r, LI, NL, SE,	FI	
JP	2000512746		T	20000926	JP	1997-541578		19970523
JP	3452068		B2	20030929				
AT	215698		Τ	20020415	AT	1997-927032		19970523
ES	2175416		Т3	20021116	ES	1997-927032		19970523
US	20020022274		A1	20020221	US	1998-194099		19981120
US	6420183		B2	20020716				
US	20020009754		A1	20020124	US	2001-966522		20010928
US	20020015969		A1	20020207	US	2001-966137		20010928
US	7063952			20060620				
US	20030092081		A1	20030515	US	2002-263607		20021003
	7138280		B2	20061121				
	20080318270		A1	20081225	IIS	2008-199317		20080827
	7615376		B2	20091110		2000 13301		20000021
	APPLN. INFO			20031110	DE	1996-19621312	70	19960528
INIONII.	. ILL DIV. THEO					1997-EP2662		19970523
						1998-194099		19981120
					US	2001-966522	A.3	20010928

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

In a procedure for quant, optical anal, of fluorescently-labeled biol, cells a cell layer is applied to a transparent carrier at the base of a reaction vessel so that it is in contact with the solution containing the fluorescent dye. The sensitivity of the determination may be significantly improved by adding to the solution a masking dye, which absorbs the exciting light for the fluorescent dye already present in the solution and/or its emitted light, and/or by applying an interlayer that is permeable to the solution but absorbs and/or reflects the exciting light or the emitted light to the cell layer at the base. The same procedure may be used to improve sensitivity in quant. optical anal. of a luminescent biol. cell layer. In the latter case the interlayer should be composed so that it possesses a high reflection factor with respect to luminescent light. Analogously, these procedural principles may also be applied in receptor studies to mask disturbing background radiation in quant. optical anal. of fluorescently- or luminescently-labeled participants in a reaction. In this case a receptor layer is placed at the base of a reaction vessel in contact with a solution (supernatant 3) in which a fluorescent or luminescent ligand has been dissolved. The sensitivity and accuracy of the determination

may

be significantly improved if a masking dye which absorbs the exciting light for the fluorescent dye and/or its emitted light or (in case of luminescent ligands) the luminescent light is added to the supernatant. An interlayer that is permeable to the solution but absorbs and/or reflects the exciting light and/or the emitted light or the luminescent light may be applied to the cell or receptor layer at the base instead of the masking dye in the solution or possibly as a supplementary measure.

masking dye in the solution or possibly as a supplementary measure.
OS.CITING REF COUNT: 14 THERE ARE 14 CAPLUS RECORDS THAT CITE THIS

RECORD (14 CITINGS)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 9 OF 13 USPATFULL on STN
ACCESSION NUMBER: 96:99016 USPATFULL <<LOGINID::20110427>>
TITLE: Skin-coloring preparation
INVENTOR(S): Kurz, Thekla, Gross-Zimmern, Germany, Federal Republic

Stossel, Sieglinde, Reinheim, Germany, Federal Republic of

Spiller, Andrea, Lemgo, Germany, Federal Republic of PATENT ASSIGNEE(S): Merck Patent Gesellschaft Mit Beschrankter Haftung, Darmstadt, Germany, Federal Republic of (non-U.S.

corporation)

NUMBER KIND DATE US 5569460 19961029 US 1994-254003 19940603 PATENT INFORMATION: 19940603 (8) APPLICATION INFO.:

NUMBER DATE PRIORITY INFORMATION: DE 1993-4318576 19930604

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

FILE SEGMENT: Granted
PRIMARY EXAMINER: Kishore, Gollamudi S.

LEGAL REPRESENTATIVE: Millen, White, Zelano, & Branigan, P.C.

NUMBER OF CLAIMS: NUMBER OF CLAIM: 1
EXEMPLARY CLAIM: 1
478

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention relates to a skin-coloring preparation, containing a hydroxycarbonyl compound which has self-tanning properties, in a cosmetologically acceptable carrier, which preparation contains at least one colorant which adheres to the skin.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L12 ANSWER 10 OF 13 USPATFULL on STN

ACCESSION NUMBER: 86:54783 USPATFULL <<LOGINID::20110427>>

TITLE: Electrooptical device having fixed translucent INVENTOR(S): Hotta, Yoshio, Atsugi, Japan
PATENT ASSIGNEE(S): Canon Kabushiki Kaisha, Tokyo, Japan (non-U.S. Electrooptical device having fixed translucent indicia

corporation)

NUMBER KIND DATE PATENT INFORMATION: US 4614407 19860930 APPLICATION INFO.: US 1983-501866 19830607 (6)

NUMBER DATE PRIORITY INFORMATION: JP 1982-102581 19820614 JP 1982-143842 19820819

DOCUMENT TYPE: Utility
FILE SEGMENT: Granted
PRIMARY EXAMINER: Corbin, John K.
ASSISTANT EXAMINER: Gallivan, Richard F.

LEGAL REPRESENTATIVE: Fitzpatrick, Cella, Harper & Scinto

NUMBER OF CLAIMS: 14 EXEMPLARY CLAIM: 9,14

NUMBER OF DRAWINGS: 4 Drawing Figure(s); 2 Drawing Page(s) LINE COUNT: 301

An electrooptical device comprises an insulating film on at least one of a pair of electrode plates, wherein the insulating film has an area defining a colored pattern.

L12 ANSWER 11 OF 13 CAPLUS COPYRIGHT 2011 ACS on STN ACCESSION NUMBER: 1971:81809 CAPLUS <<LOGINID::20110427>>

74:81809 DOCUMENT NUMBER:

ORIGINAL REFERENCE NO.: 74:13243a,13246a

TITLE: Photoconducting recording materials

INVENTOR(S): Tavernier, Bernard H.; Vanheertum, Johannes J.

PATENT ASSIGNEE(S): Gevaert-Agfa N. V.
SOURCE: Belg., 10 pp.
CODEN: BEXXAL

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PRIORITY APPLN. INFO.: GB 19681112
AB To mask the yellow-to-brown color of PbO as photoconductor it is

overcoated with a white pigment (TiO2, SiO2, BaSO4) layer containing <40% of a binder, which is transparent to x-rays, but reflects visible light, so that the materials may be charged, x-ray exposed, and processed in subdued daylight or artificial light. Thus, a dispersion of 50 g of a com. yellow PbO in 100 g PhMe containing 0.5 g monobutyl phosphate was ballmilled with 8-mm diameter ceramic soherules, and after addition of 15 ml of a 50% PhMe

solution

of styrens-modified alkyd resin (Alkydal V-15), for an addnl. 128 hr. The dispersion was coated on Al-clad paper to give a 60- $\mu$  Layer of 150 g PbO/m2 and overcoated with 20 g/m2 of TiO2 (0.5-10 $\mu$  particle size) in the form of a dispersion of TiO2 25 g, poly(vinyl acetate) 50 g, and monobutyl phosphate 250 mg in PhMe 100 ml. A Shellsol T developer containing carbon black and Alkydal L-67 resin yielded black radiographs on a brilliant white background.

L12 ANSWER 12 OF 13 CAPLUS COPYRIGHT 2011 ACS on STN

ACCESSION NUMBER: 1971:59406 CAPLUS <<LOGINID::20110427>>

DOCUMENT NUMBER: 74:59406

ORIGINAL REFERENCE NO.: 74:9561a,9564a

TITLE: Photoconductive recording material

TITAID DAME

INVENTOR(S): Tavernier, Bernard H.; De Meyer, Alfons J.;

Vanheertum, Johannes J.

PATENT ASSIGNEE(S): Gevaert-Agfa N. V. SOURCE: Belg., 13 pp.

CODEN: BEXXAL DOCUMENT TYPE: Patent

LANGUAGE: French FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAIENI NO.	KIND	DAIE	APPLICATION	NO.	DAIE
	BE 740943		19700429	BE		19691029
	DE 1954633			DE		
	FR 2023143			FR		
	GB 1280023			GB		
	US 3642470		19720215	US		19691112
PRIOR	RITY APPLN. INFO.:			GB		19681112
AB	To improve the color	of ele	ctrophotog.	records made	with x-ray	s or visible
	radiation using teti					
	PbO (U.S. 3,266,932)	CA 65:	13068) the	layer, pref	erably poly	/(vinyl

radiation using tetragonal (U.S. 3,008,825; CA 56: 5572b) or orthorhombic PbO (U.S. 3,266,932; CA 66: 13068) the layer, preferably poly(vinyl acetate) with 50-90% red or brownish PbO is treated prior to or after the exposure with a >25% solution of a NH4, alkali metal, or alkaline earth metal

halide. The effect of CaCl2 is described as 9 PbO + 3 CaCl2 + 9 H2O  $\rightarrow$  2(3 Pb(OH)2.PbCl2) + PbCl2 + 3 Ca(OH)2. The charge acceptance is not markedly changed by the treatment, and the light-sensitivity lowered somewhat. Thus, the brownish color of a layer of 80% tetragonal PbO in poly(vinyl acetate) on an Al plate is masked by treatment for 15 sec at 80° with 25 aqueous CaCl2. The x-ray-exposed material yields a black image on a brilliant white background upon processing with a liquid developer containing carbon black as toner.

L12 ANSWER 13 OF 13 CAPLUS COPYRIGHT 2011 ACS on STN
ACCESSION NUMBER: 1919:7380 CAPLUS <<LOGINID::20110427>>
DOCUMENT NUMBER: 13:7380
ORIGINAL REFERENCE NO.: 13:1396c-g
TITLE: Acid dyes containing chromium
PATENT ASSIGNEE(S): Soc. Anon. Pour L'ind. Chim. A Bale

SOURCE: Additions to 77,662 (C. A. 13, 191).
DOCUMENT TYPE: Patent

LANGUAGE: Unavailable

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

CH 78615 19180816 CH

Swiss, 78,615 to 78,625, inclusive, all dated Aug. 16, 1918. In the manufacture of Cr-containing acid dyes, the azo dyes, reactive to metals, obtained by any of the methods specified below, are treated with Cr208 or its salts until the reaction is complete. The initial materials specified for the preparation of the azo dyes are (a) diazotized 1-amino-2-hydroxynaphthalene-4-sulfonic acid and  $\alpha$ -naphthol, (b) diazotized 1-amino-2-hydroxynaphthalene-4-sulfonic acid and 1-phenyl-3-methyl-5-pyrazolone, (c) diazotized 4-nitro-2-amino-1-hydroxybenzene and 1-hydroxy-naphthalene-5-sulfonic acid, (d) diazotized 4,6-dinitro-2-amino-1-hydroxybenzene and 1,8-aminonaphthol-2,4-disulfonic acid, (e) diazotized 4-chloro-2-amino-1-hydroxy-benzene-6-carboxylic acid and 1,8-aminonaphthol-2,4-disulfonic acid, (f) diazotized 4-chloro-2-amino-1-hydroxybenzene-6-sulfonic acid and 1,8-aminonaphthol-,3,6-disulfonic acid, (g) diazotized 4-chloro-2-amino-1-hydroxybenzene and benzoylacetic-0-carboxylic acid, (h) diazotized 1-amino-2-hydroxynaphthalene-4-sulfonic acid and 3-hydroxy-1-thionaphthene, (i) diazotized anthranilic acid and benzoyl-2-amino-5-naphthol-7-sulfonic acid, (j) diazotized anthranilic acid and phthaloy1-2-amino-5-naphthol-7-sulfonic acid, (k) diazotized 1-amino-2-hydroxynaphthalene-4-sulfonic acid and diketohydrindene. The dye products are soluble in H2O, contain Cr in masked form (i. e., the Cr cannot be precipitated from aqueous solution by Na2CO3, NaOH, or

NH4OH), and dye
animal fibers from acid bath in fast colors, i. e., resp., blue, blue-red,
yellowish brown-black, yellowish green, brilliant
blue, clear green-blue, clear yellowish green, greenish blue,
bordeaux-red, and prune, shades.

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(FILE 'HOME' ENTERED AT 14:52:53 ON 27 APR 2011)

FILE 'CAPLUS, MEDLINE, BIOSIS, BIOTECHNO, COMPENDEX, ANABSTR, CERAB, METADEX, USPATFULL' ENTERED AT 14:54:25 ON 27 APR 2011

122 SEA FILE-MYFE SPE-ON ABB-ON PLU-ON (FLUORESCEN? (3A) DYE)

L2	122	(P) (MASKING (3A) DYE) DUP REM L1 (0 DUPLICATES REMOVED) D L2 1-20 TI
* 0		D L1 21-122 TI D L1 100-122 IBIB ABS
L3	ь	SEA FILE-MFE SPE-ON ABB=ON PLU-ON (FLUORESCEN? (3A) DYE) (P) (MASKING (3A) BACKGROUND (5A) DYE) D L3 1-6 TI
L4	4	SEA FILE=MFE SPE=ON ABB=ON PLU=ON (FLUORESCEN? (3A) DYE) (P) (SECOND (3A) DYE) (P) (REDUC? (3A) BACKGROUND) D L4 TI D L4 1-4 TI D L4 1-4 TI D L4 1-1 IBIB ABS E REYNOLDS C PATRICK/AU
L5	12	SEA FILE=MFE SPE=ON ABB=ON PLU=ON "REYNOLDS C PATRICK"/AU AND (DYE)
L6	7	DUP REM L5 (5 DUPLICATES REMOVED) D L6 1-7 IBIB TI E FRGALA TOMAS/AU
L7	7	SEA FILE=MFE SPE=ON ABB=ON PLU=ON "FRGALA TOMAS"/AU AND DYE
L8	4	DUP REM L7 (3 DUPLICATES REMOVED) D L8 1-4
L9	29	SEA FILE=MFE SPE=ON ABB=ON PLU=ON (FLUORESCENT (3A) DYE) (P) (BRILLIANT (3A) BLACK)
L10	29	DUP REM L9 (0 DUPLICATES REMOVED) D L10 1-29 TT
L11	13	SEA FILE=MFE SPE=ON ABB=ON PLU=ON (BRILLIANT (3A) BLACK) (P) (MASK?)